=> d ibib ab hitstr 1-11

4/28/63

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L4 ANSWER 1 OF 11
ACCESSION NUMBER:
TITLE:

Process for preparing17alpha-acetoxy-llbeta-{4-n, n(disethylamino)phenyl}-2l-methoxy-19-norpregna-4,9-diene -3,20-dione, intermediates useful in the process, and processes for preparing such intermediates

INVENTOR(S):

Kim, Hyun Koo, Bethesda, MD, UNITED STATES
Rao, Pemmaraju N, San Antonio, TX, UNITED STATES
Simmons, Anne Marie, San Antonio, TX, UNITED STATES
                                                                                                                                                       NUMBER
                                                                                                                            US 2003060646 Al 20030327

US 2002-169139 Al 20020627 (10)

WO 2000-US35479 20001229

Utility

APPLICATION

LEVDIG VOIT & MAYER, LTD, 700 THIRTEENTH ST. NW, SUITE

300, WASHINGTON, DC, 20005-3960
     PATENT INFORMATION:
APPLICATION INFO.:
     DOCUMENT TYPE:
FILE SEGMENT:
LEGAL REPRESENTATIVE:
LEGAL REPRESENTATIVE: LEYDIG VOIT & MAYER, LTD, 700 THIRTEENTH ST. NW, SUITE 300, WASHINGTON, DC, 20005-3960

NUMBER OF CLAIMS: 44

EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 2 Drawing Page(s)

LINE COUNT: 1219

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A compound having general formula (I) in which R.sup.l is a member selected from the group consisting of --OCH.sub.3, --SCH.sub.3, --N(CH.sub.3).sub.2, --NHCH.sub.3, --CHO, --COCH.sub.3 and --CHOKCH.sub.3 is R.sup.2 is a member selected from the group consisting of halogen, alkyl, acyl, hydroxy, alkoxy, acyloxy, alkyl carbonate, cypionyloxy, S-alkyl and S-acyl; R.sup.3 is a member selected from the group consisting of flydrogen and alkyl; and X is a member selected from the group consisting of --O and --N-OR.sup.5, wherein R.sup.5 is a member selected from the group consisting of flydrogen and alkyl. In addition to providing the compounds of formula (I), the present invention provides methods wherein the compounds of formula (I) are advantageously used, inter alia. to antagonize endogenous progesterone; to induce enness; to treat endometriosis; to treat dysmenorrhea; to treat endocrine hormone-dependent tumors; to treat dysmenorrhea; to treat endocertine hormone-dependent tumors; to induce labor; and for contraception. ##STRI##

[1 19841-31-2P (process for the prepn. of 17.alpha-acetoxy-11.beta.-[4-N.N-(dimethylaming)leheiu]-21-methoxy-19-noprepagena-4-9-diene-3 #D-diene-3
                             (process for the prepn. of 17.alpha.-acetoxy-11.beta.-[4-N,Ny
  (dimethylamino)phenyl]-21-methoxy-19-norpregna-4,9-diene-3,30-dione,
  intermediates useful in the process, and processes for prefig. such
  intermediates)
198414-31-2 USPATFULL
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-
  (dimethylamino)phenyl]-21-methoxy-, (11.beta.)- (9CI) (CA INDEX NAME
                                                                                                                                                                                                                                                                                                                                 (CA INDEX NAME)
                                       Absolute stereochemistry.
  L4 ANSWER 2 OF 11
ACCESSION NUMBER:
TITLE:

USPATFULL
2002:301659 USPATFULL
1mplantation rates after in vitro fertilization, and treatment of intirtility and early pregnancy loss with a nitric oxide gonor or substrate alone or in combination with projecterone, and a method for contraception with nitric oxide inhibitors in combination yith antiprogesting or other agents Chwalisz, Kfzysztof, Berlin, GERMANY, FEDERAL REPUBLIC OF
                                                                                                                                                                                  Robert E., Friendswood, TX, UNITED STATES
                                                                                                                           NUMBER KIND DATE

US 2002169205 A1 20021114
US 2002-43232 A1 20020114 (10)
Dividion of Sec. No. US 1998-162446, filed on 29 Sep
1998, PENDING
ULSPILICY
ABBLICATION
MILLEN, WHITE, ZELANO & BRANIGAN, P.C., 2200 CLARENDON
BLVD., SUITE 1400, ARLINGTON, VA, 22201
47
     PATENT INFORMATION:
APPLICATION INFO.:
RELATED APPLN. INFO.:
     DOCUMENT TYPE:
FILE SEGMENT:
LEGAL REPRESENTATIVE:
   NUMBER OF CLAIMS:
EXEMPLARY CLAIM:
NUMBER OF DRAWINGS:
LINE COUNT:
                                                                                                                                        Drawing Page(s)
  LINE COUNT:

AB A method is provided for the improvement of implantation rates and/or pregnancy takes in a female mammal, comprising administering to a female mammal in show pregnancy is desired an effective amount of
                                         (a) a nifric oxide synthase substrate, a nitric oxide donor, or both, optionally in combination with
                                 (c) optionally, in further combination with an estrogen. A method is also provided for fertility control for a female mammal, comprising administering to a female mammal in whom pregnancy is not desired and at risk for becoming pregnant an effective amount of nitric oxide synthase inhibitor in combination with an antiprogestin. Pharmaceutical fompositions are also provided. (774-99-4, CDB 2914 (antiprogestin, method for contraception with nitric oxide inhibitors in combination with antiprogestins or other agents) 126784-99-4 USPATPULL 19-Norpregna-4,9-diene-3,20-diene, 17-(acetyloxy)-11-(4-(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)
                                       Absolute stereochemistry.
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ANSWER 1 OF 11 USPATFULL ANSWER 2 OF 11 USPATFULL (Continued)

L4 ANSWER 3 OF 11 USPATFULL

ACCESSION NUMBER: 2002:43584 USPATFULL

21-SUBSTITUTED PROGESTERONE DERIVATIVES AS NEW ANTIPROGESTATIONAL AGENTS

INVENTOR(S): KIM, HYUN K., BETHESDA, MD, UNITED STATES

RAO, PEPHARAJU N., SAN ANTONIO, TX, UNITED STATES

RAO, PEPHARAJU N., SAN ANTONIO, TX, UNITED STATES

ACOSTA, CARMIE K., SAN ANTONIO, TX, UNITED STATES

ACOSTA, CARMIE K., SAN ANTONIO, TX, UNITED STATES

NUMBER KIND DATE US 2002025951 US 1999-180132 WO 1997-US7373 Utility APPLICATION 20020228 19990524 19970430 PATENT INFORMATION: APPLICATION INFO.: (9)

DOCUMENT TYPE: EUGENIA GARRETT WACKOWSKI, TOWNSEND AND TOWNSEND AND CREW, TWO EMBARCADERO CENTER, 8TH FLOOR, SAN FRANCISCO, CA, 94111 FILE SEGMENT: LEGAL REPRESENTATIVE:

NUMBER OF CLAIMS: EXEMPLARY CLAIM: NUMBER OF DRAWINGS: LINE COUNT:

NUMBER OF DRAWINGS: 3 Drawing Page(s)
LINE COUNT: 2185
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AB A compound having the general formula: ##STR1##

in which: R.sup.1 is a member selected from the group consisting of --OCH.sub.3, --SCH.sub.3, --N(CH.sub.3).sub.2, --NHCH.sub.3, --CHO, --COCH.sub.3 and --CHONCH.sub.3, sup.2 is a member selected from the group consisting of halogen, alkyl, acyl, hydroxy, alkoys, acyloxy, alkyl carbonate, cypionyloxy, S-alkyl and S-acyl: R.sup.3 is a member selected from the group consisting of alkyl, hydroxy, alkoys and acyloxy: R.sup.4 is a member selected from the group consisting of hydrogen and alkyl: and X is a member selected from the group consisting of .sub..dbd.0 and .sub..dbd.N-OR.sup.5, wherein R.sup.5 is a member selected from the group consisting of .sub..dbd.0 and .sub..dbd.N-OR.sup.5, wherein R.sup.5 is a member selected from the group consisting of hydrogen and alkyl.

In addition to providing the compounds of Formula I, the present invention provides methods wherein the compounds of Formula I are advantageously used, inter alia, to antagonize endogenous progesterone; to induce menses; to treat endometriosis; to treat dysmenorches; to treat endocrine hormone-dependent tumors; to treat uterine fibroids; to inhibit uterine endometrial proliferation; to induce labor; and for contraception.

contraception. IT 198414-07-2P 198414-31-2P

(prepn. of progesterone derivs. as antiprogestational agents)
198414-07-2 USATFULL
198414-07-2 USATFULL
19-Norpregna-4.9-diene-3,20-dione, 17,21-bis(acetyloxy)-11-(4(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 3 OF 11 USPATFULL (Continued)
198414-05-0 USPATFULL
198414-05-0 USPATFULL
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-21-chloro-11-(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

198414-11-8 USPATFULL 19-Norpregna-4,9-diene-3,20-dione, 17-(acety) (dimethylamino)phenyl]-, (11.beta.)- (9CI) oxy)-21-(acetylthio)-11-[4-(CA INDEX NAME)

Absolute stereochemistry.

198414-22-1 USPATFULL
Estra-4,9-dien-3-one, 17-(acctyloxy)-11-[4-(dimethylamino)phenyl]-17-(1oxopropyl)-, (11.beta.,17.alpha.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

198414-33-4 USPATFULL
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-21-(3-cyclopentyl-1-oxopropoxy)-11-(4-(dimethylamino)phenyl)-, (11.beta.)- (9CI) (CA INDEX

ANSWER 3 OF 11 USPATFULL

198414-31-2 USPATFULL 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4-(dimethylamino)phenyl]-21-methoxy-, (11.beta.)-_[9CI) (CA INDEX NAME) Absolute stereochemistry.

IT 198414-03-8P 198414-05-0P 198414-11-8P 198414-22-1P 198414-33-4P 198414-34-5P 198414-39-0P 198414-43-6P (prepn. of progesterone derivs. as antiprogestational agents)
RN 198414-03-8 (USPATFULL CN 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-21-fluoro-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 3 OF 11 USPATFULL NAME) (Continued)

Absolute stereochemistry.

198414-34-5 USPATFULL 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-21-hydroxy-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

198414-39-0 USPATFULL
19-Norpreyna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-21-ethoxy-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

198414-43-6 USPATFULL
19-Norpregna-4,9-diene-3,20-diene, 17-(acetyloxy)-21-bromo-11-[4-(diesthylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

ANSWER 3 OF 11 USPATFULL

Absolute stereochemistry.

198414-40-3P 198414-41-4P
(prepn. of progesterone derivs. as antiprogestational agents)
198414-40-3 USPATFULL
19-Norpregna-4,9-diene-3,20-dione, 17,21-bis(acetyloxy)-11-{4(dimethylamino)phenyl}-, 3-oxime, (3E,11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

198414-41-4 USPATFULL
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylmaino)phenyl]-21-methoxy-, 3-oxime, (11.beta.}- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry unknown.

L4 ANSWER 4 OF 11 USPATFULL
ACCESSION NUMBER: 2000:34586 USPATFULL
TITLE: Implantation rates after in vitro fertilization, treatment of infertility and early pregnancy loss with a nitric oxide donor alone or in combination with progesterone, and a method for contraception with nitric oxide inhibitors

INVENTOR(S): Chwalisz, Krzysztof, Berlin, Germany, Federal Republic of

of Garfield, Robert E., Friendswood, TX, Unyted States Schering Aktiengesellschaft, Berlin, Gefanny, Federal Republic of (non-U.S. corporation) The Board of Regents, Univ. of Texas System, Austin, TX, United States (U.S. corporation)

PATENT ASSIGNEE (S):

NUMBER KIND DATE

US 6040340 20000321 US 1996-646518 19960507 (8) Utility Granted MacHillan, Keith D. Millen, White, Zelano & Branigan, P.C. 27 PATENT INFORMATION: US 6040340 20000321
APPLICATION INFO: US 1996-646518 1996507 (8)
DOCUMENT TYPE: Utility
FILE SEGMENT: Granted
PRIMARY EXAMINER: MacMillan, Keith D.
LEGAL REPRESENTATIVE: Millen, White, Zelano & Branigan, P.C.
NUMBER OF CLAIMS: 27
EXEMPLANY CLAIM: 1
NUMBER OF DRAWINGS: 6 Drawing Figure(s): 6 Drawing Page(s)
LINE COUNT: 756
CAS INDEXING IS AVAILABLE FOR THIS PATENT
AB A method is provided for the improvement of implantation rates and/or pregnancy rates in a female mammal, comprising administering to a female mammal in whom pregnancy is deared an effective amount of

(a) a nitric oxide synthase substrate, a nitric oxide donor, or both, optionally in combination with

(b) a progestin, and,

(c) optionally, in further combination with an estrogen.

A method is also provided for fertility control for a female mammal, comprising administeding to a female mammal in whom pregnancy is not desired and at risk for becoming pregnant an effective amount of nitric oxide synthase inhibitor in combination with an antiprogestin. Pharmaceutical compositions are also provided.

IT 120784-99-4, CD23914

(fertility control using a nitric oxide synthase inhibitor in combination with an antiprogestin)

RN 126784-99-4 USPATFULL

(31-Norpregnay4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

ANSWER 3 OF 11 USPATFULL (Continued)

ANSWER 4 OF 11 USPATFULL (Continued)

L4 ANSWER 5 OF 11 USPATFULL ACCESSION NUMBER: 2000 PAFFULL
2000:12791 USPATFULL
20-keto-11.beta.-arylsteroids and their derivatives having agenist or antagonist hormonal properties Cook, C. Edgar, Staunton, VA, United States Kepler, John A., Raleigh, NC, United States Zhang, Ping-sheng, Hillbrae, CA, United States Lee, Yue-wei, Chapel Hill, NC, United States Tallent, C. Ray, Raleigh, NC, United States Tallent, C. Ray, Raleigh, NC, United States, NC, United States Tallent, C. Ray, Raleigh, NC, United States Tallent, C. Ray, Raleigh, NC, United States Newscarch Triangle Park, NC, United States (U.S. corporation) INVENTOR(S): PATENT ASSIGNEE(S): NUMBER KIND DATE NUMBER KIND DATE

PATENT INFORMATION: US 6020328 20000201
APPLICATION INFO: US 1998-35949 19980306 (9)
DOCUMENT TYPE: Utility
FILE SECRET: Granted
PRIMARY EXAMINER: Beds, Jose' G.
ASSISTANT EXAMINER: Beds, Jose' G.
ASSISTANT EXAMINER: Beds, Spivak, McCleiland, Maier & Neustadt, P.C.
NUMBER OF CLAIMS: 9
EXEMPLARY CLAIM: 1 1
STAMINER OF DRAWINGS: 5 Drawing Figure(s); 10 Drawing Page(s)
LINE COUNT: 2599
CAS INDEXING STAVALLEBLE FOR THIS PATENT.
AB The invention is directed to 20-keto-11.beta.-arylsteroids of formula I:
##STRIFF wherein R.sup.1, R.sup.6, R.sup.7, R.sup.9, R.sup.12 and X are as defined by the specification. The compounds exhibit progestational and antiprogestational activities.

IT 240805-97-4P 240806-96-8P
240806-03-5P 240806-98-5P 240805-99-5P
240805-97-4P 240805-98-5P 240805-98-5P
240805-97-4P 240805-98-5P 240805-98-5P
340805-97-4P 240805-98-5P 240805-98-5P
240805-97-4P 240805-98-5P 240805-98-5P
340805-97-4P 240805-98-5P 240805-98-6P
340805-97-4P 240805-98-5P 240805-98-5P
340805-98-5P 240805-98-5P
3408 Absolute stereochemistry. 240805-98-5 USPATFULL
19-Norpregna-4,9-diene-3,20-dione, 11-[4-(dimethylamino)phenyl]-17[(phenylacetyl)oxyl-, (11.beta.)- (9CI) (CA_XNOEX NAME) ANSWER 5 OF 11 USPATFULL (Continued) Absolute stereochemistry

L4 ANSWER 5 OF 11 USPATFULL Absolute stereochemistry 240805-99-6 USPATFULL 19-Norpregna-4,9-diene-3,20-dione, 17-(benzoyloxy)-11-[4-(dimethylamino)pheny1]-, (11.beta.)- (9CI) (CA INDEX NAME) Absolute stereochemistry. Me₂N 240806-03-5 USPATFULL 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4-methoxyphenyl)-, (11.beta.)- (9CI) (CA INDEX NAME) Absolute stereochemistry. 240806-06-8 USPATFULL 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)-3-fluorophenyl]-, (11.beta.)- (9CI) (CA INDEX NAME) L4 ANSWER 6 OF 11 USPATFULL

ACCESSION NUMBER:

TITLE: Hethod for preparing 17.alpha.-acetoxy-11.beta.-(4-N, N-dimethylaminophyl)-19-Norpregna-4,9-diene-3, 20-dione, intermediates useful in the method, and methods for the preparation of such intermediates Xim, Hyun K., Betheada, MD, United States Rao, Pemmaraju Narasinha, San Antonio, TX, United States Burdett, Jr., James E., Somerset, TX, United States Acosta, Carmie K., San Antonio, TX, United States The United States of America as represented by the Department of Health and Human Services, Vashington, DC, United States (U.S. corporation) PATENT INFORMATION: US 5929262 19990727
APPLICATION INFO: US 1995-413755 19950330 (8)
DOCUMENT TYPE: UTILITY
FILE SECMENT: Granted
PRIMARY EXAMINER: Bedio, Barbara
Legal REPRESENTATIVE: Leydig, Voit & Mayer
NUMBER OF CLAIMS: 19
EXEMPLARY CLAIM: 1
NUMBER OF DRAWINGS: 2 Drawing Figure(s); 2 Drawing Page(s)
LINE COUNT: 777
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AB Methods for the preparation of the 19-norprogesterone of formula I ##STRI## and its intermediates, in crystalline and amorphous forms are disclosed. The process is performed by (1) protecting the hydroxyl group of a compound of formula II ##STR2## (2) reacting the protected compound with an alkali or alkaline earth metal anion radical, (3) hydrolyzing the resulting compound, (4) ketalizing the carbonyl groups, (5) epoxidizing the compound, (6) opening the epoxide ring and introducing an N,N,dimethylamino-phenyl functional group into the axial position of C. sub.11, (7) deketalizing and dehydrating the resulting compound, and (8) acetylating to provide 17.alpha.-acetoxy-11.beta.-(4-N,N-dimethylaminophenyl)-19-norpregna-4,9-diene-3,20-dione (I).

IT 126784-99-4 USPAFFULL

RN 126784-99-4 USPAFFULL

RN 126784-99-4 USPAFFULL

ANSWER 7 OF 11 USPATFULL L4 (Continued) OHO

L4 ANSWER 7 OF 11 USPATFULL

ACCESSION NUMBER:

TITLE:

11 .beta.-phenyl-gonanes, their manufacture and pharmaceutical preparations containing them

Neef, Gunter, Berlin, Germany, Federal Republic of Elger, Walter, Berlin, Germany, Federal Republic of Henderson, David, Berlin, Germany, Federal Republic of Otto, Eckard, Berlin, Germany, Federal Republic of Rohde, Ralph, Berlin, Germany, Federal Republic of Schering Aktiengesellschaft, Berlin and Bergkamen, Germany, Federal Republic of Schering Aktiengesellschaft, Berlin and Bergkamen, Germany, Federal Republic of Non-U.S. corporation) INVENTOR(S): PATENT ASSIGNEE(S): PATENT INFORMATION: APPLICATION INFO.: US 5089635 US 1986-827050 19920218 19860207 NUMBER DATE DE 1985-3504421 1985
DE 1985-3527517 1985
Utility
Granted
Killos, Paul J.
Millen, White & Zelano PRIORITY INFORMATION: 19850207 19850729 DOCUMENT TYPE: DOCUMENT TYPE: Utility
FILE SEGMENT: Granted
PRIMARY EXAMINER: Killos, Paul J.
LEGAL REPRESENTATIVE
MIMBER OF CLAIMS: 45
EXEMPLARY CLAIM: 1
LINE COUNT: 128
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AB 13-alkyl-11.beta.-phenyl-gonanes of general formula I ##STRI## wherein A
and B together stand for an oxygen atom, a CH.sub.2 group or a second
bond between carbon atoms 9 and 10, X is an oxygen atom or the hydroxyimino grouping N. about. OH, R.sub.l is a straight-chained or branched, saturated or unsaturated alkyl radical with up to 8 carbon atoms, which contains the grouping 45578249 with X as described above, R.sub.2 is a methyl or ethyl radical in the .alpha. or .beta. position, R.sub.9, R.sub.10, R.sub.11 and R.sub.12 each stand for a hydrogen atom, a hydroxy, alkyl, alkoxy or acyloxy group with 1 to 4 carbon atoms respectively or a halogen atom and R.sub.3 and R.sub.4 have a variety of meanings, have antigestagenic and antiglucocorticoid effects.

17 105114-79-2P (prepn. of, as antigestagen and antiglucocorticoid)
105114-79-2 USPATFULL Benzaldehyde, 4-[(11.beta.,13.alpha.)-17-(acetyloxy)-3,20-dioxo-19-norpregna-4,9-dien-11-yl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

4 ANSWER 8 OF 11 USPATFULL
CCESSION NUMBER: 91:102214 USPATFULL
11 .beta. -substituted progesterone analogs
Cook, C. Edgar, Durham, NC, United States
Wani, Hansukh C., Durham, NC, United States
Lee, Yun W., Chapel Hill, NC, United States
Reel, Jerry R., Cary, NC, United States
Rector, Douglas, Mobile, AL, United States
ATENT ASSIGNEE(S): Research Triangle Park,
NC, United States (U.S. corporation) TITLE: INVENTOR(S):

PATENT ASSIGNEE(S):

NUMBER KIND DATE

US 5073548 19911217
US 1990-504129 19900403 (7)
Division of Ser. No. US 1988-210503, filed on 23 Jun 1988, now patented, Pat. No. US 4954490
Utility
Granted
Shab. Nukund J. PATENT INFORMATION:

APPLICATION INFO.: RELATED APPLN. INFO.:

DOCUMENT TYPE:

DOCUMENT TYPE:
FILE SEGMENT:
PRIMARY EXAMINER:
ASSISTANT EXAMINER:
LEGAL REPRESENTATIVE:
NUMBER OF CLAIMS:
NUMBER OF DRAWINGS:
LIME COUNT.

Shah, Mukund J. Ward, E. C. Oblon, Spivak, McClelland, Maier & Neustadt

2 Drawing Figure(s); 2 Drawing Page(s)

NORDER OF DOMESTICS:
LINE COUNT:

LINE COUNT:

AB A 11.beta.acyl-19-norprogesterone steroid of the formula: ##STR1##

Wherein (1) R.sup.1 is H. C.sub.1-4 alkyl, C.sub.2-4 alkenyl, C.sub.2-4

alkynyl, OH. OC (O)CH.sub.3, or OC (O)R.sup.5, b. wherein R.sup.5 is

C.sub.2-8 alkyl, C.sub.2-8 alkenyl, C.sub.2-8 alkynyl or acyl, R.sub.2

is H. R.sup.3 is H. C.sub.1-4 alkyl, C.sub.2-8 alkynyl or c.sub.2-4

alkynyl, R.sup.4 is H. CH.sub.3, F or Cl, R.sup.6 is H. (CH.sub.3).sub.2

N, CH.sub.3 O, CH.sub.3 CO, CH.sub.3 S, CH.sub.3 SO, CH.sub.3 SO.sub.2,

and X is O or NOCH.sub.3; or and X is O or NOCH.sub.3 ; or

(ii) R.sup.1 and R.sup.2 taken together are a carbon-carbon bond and R.sup.3, R.sup.4, R.sup.6 and X are as defined above; or

(iii) R.sup.1 and R.sup.3 taken together are --CH.sub.2 -- or --N.dbd.N--CH.sub.2 --, R.sup.2 is H and R.sup.4, R.sup.6 and X are as defined above; or

(iv) R.sup.2 and R.sup.3 taken together are .dbd.CH.sub.2 and R.sup.1 , R.sup.4, R.sup.6 and X are as defined above. IT 12659-26-4P 12690-23-7P 126784-99-4P

(CA INDEX NAME)

ANSWER 8 OF 11 USPATFULL

126690-29-7 USPATFULL
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4-acetylphenyl)-,
(11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereoghemistry.

126784-99-4 USPATFULL 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-{4-(dimethylamino)phenyl}-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 9 OF 11 USPATFULL

-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4-acetylphenyl)-, (11.beta.)- (9C1) (CA INDEX NAME)

Absolute stereochemistry.

126784-99-4 USPATFULL 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

stereochemistry. Absolute Meak

L4 ANSWER ACCESSION NU TITLE: INVENTOR(S):

4 ANSWER 9 OF 11 USPATFULL
CCESSION NUMBER: 90:69718 USPATFULL
11 .beta.-substituted progesterone analogs
Cook, C. Edgar, Durham, NC, United States
Wani, Mansukh C., Research Triangle Park, NC, United
States
Lee, Y.-W, Chapel Hill, NC, United States
Reel, Jerry R., Delmar, NY, United States
Rector, Douglas, Raleigh, NC, United States
Research Triangle Park,
NC, United States (U.S. corporation)

PATENT ASSIGNEE(S):

KIND DATE NUMBER

NUMBER KIND DATE

PATENT INFORMATION: US 4954490 19900904

APPLICATION INFO: US 4954490 19800623 (7)

DOCUMENT TYPE: Utility
FILE SEGMENT: Granted

PRIMARY EXAMINE: Lipovsky, Joseph A.

LEGAL REPASESNTATIVE: Oblon, Spivak, McClelland, Maier & Neustadt

NUMBER OF CLAIMS: 31

EXDPLARY CLAIM: 1 4 Drawing Figure(s): 1 Drawing Page(s)

LINE COUNT: 1259

AB A 11.beta.-aryl-19-norprogesterone steroid of the formula: ##STR1##

wherein (1) R. sup. 1 is H, C. sub. 1-4 alkyl, C. sub. 2-4 alkenyl, C. sub. 2-4

alkynyl, OH, OC(0)CH. sub. 3, or OC(0)R. sup. 5, wherein R. sup. 5 is

C. sub. 2-8 alkyly, C. sub. 2-8 alkenyl, C. sub. 2-4

alkynyl, R. sup. 4 is H, C. H, sub. 3; F or Cl, R. sup. 6 is H, C.H. sub. 3, or Cl, Sub. 6 alkenyl or C. sub. 2-4

alkynyl, R. sup. 4 is H, C. sub. 1-4 alkyl, C. sub. 2-4 alkenyl, C. sub. 2-4

alkynyl, R. sup. 4 is H, C. H, sub. 3; F or Cl, R. sup. 6 is H, C.H. sub. 3; or Cl, Sub. 6 alkenyl or C. sub. 2-4

and X is O or NOCH. sub. 3; or Cl sub. 3 so, CH. sub. 3 So, CH. sub. 3 So, Sub. 2,

and X is O or NOCH. sub. 3; or

(ii) R.sup.1 and R.sup.2 taken together are a carbon-carbon bond and R.sup.3, R.sup.4, R.sup.6 and X are as defined above; or

(iii) R.sup.1 and R.sup.3 taken together are --CH.sub.2 -- or --N.dbd.N--CH.sub.2 --, R.sup.2 is H and R.sup.4, R.sup.6 and X are as defined above; or

(iv) R.sup.2 and R.sup.3 taken together are .dbd.CH.sub.2 and R.sup.1,
R.sup.4, R.sup.6 and X are as defined above.

IT 12689-26-4P 126890-29-7P 126784-99-9

(prepn. of, as antiglucocorticoid and/or (anti)progestogen)

RN 126690-26-4 USPATFULL

CN 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4
(dimethylamino)phenyl]-6-methyl-, (6.alpha.,11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L4 ANSWER 10 OF 11 USPATFULL
ACCESSION NUMBER: 90.23597 USPATFULL
TITLE: Novel 11 .beta.-alkynylphenyl-10-nor-steroids
Teutsch, Jean-Georges, Pantin, France
Kilch, Hichel, Villemomble, France
PATENT ASSIGNEE(s): ROUSSel Uclaf, Paris, France (non-U.S. corporation)

NUMBER KIND DATE

US 4912097 US 1987-44958 PATENT INFORMATION: APPLICATION INFO.; 19900327 19870430 (7)

NUMBER DATE FR 1986-6517

PRIORITY INFORMATION: FR 1986-5517 19860506

DOCUMENT TYPE: Utility
FILE SEGMENT: Branch Berch, Mark L.

LEGAL REPRESENTATIVE: Bierman & Muserlian

NUMBER OF CLAIMS: 21

EXEMPLARY CLAIM: 1,9

LINE COUNT: 2174

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Novel 11.beta.—alkynylphenyl-19-nor-steroids of the formula ##STRl## wherein R. sub.1 is alkynyl of 2 to 8 carbon atoms optionally substituted with at least one member of the group consisting of --OH halogen, trialkylsilyl of 1 to 6 alkyl carbon atoms and alkylthio of 1 to 6 carbon atoms and dialkylamino of 1/to 6 alkyl carbon atoms having remarkably antiprogestosmietic and/antiplucocorticoidal activity.

IT 116421-73-99 116421-74-09

(prepn. of, as drug)

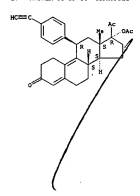
(prepn. of, as drug)
116421-73-9 USPATFULL
19-Norprepan-4, 9-diene-3, 20-diene, 17- (acetyloxy)-11-[4-(1-propyny1)pheny1]-, (11.bet4.)- (9CI) (CA INDEX NAME)

Absolute stereochemist

116421-74-0 USPATFULL

9-Norp/egna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4-ethynylphenyl)-, (11-6eta.)- (9CI) (CA INDEX NAME)

L4 ANSWER 10 OF 11 USPATFULL (Contin



RN 96286-50-6 USPATFULL
(N 18,49-Dinorpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4-(dimethylamino)phenyl)-13-ethyl-, (11.beta.,13.alpha.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

LI ANSWER 11 OF 11 USPATFULL

ACCESSION NUMBER: 88:69168 USPATFULL

13.alpha.-alkyl-gonanes, their production, and pharmaceutical preparations containing same Neef, Gunter, Berlin, Germany, Federal Republic of Beler, Sybille, Berlin, Germany, Federal Republic of Elger, Walter, Berlin, Germany, Federal Republic of Elger, Walter, Berlin, Germany, Federal Republic of Henderson, David, Berlin, Germany, Federal Republic of Schering Aktiengesellschaft, Berlin and Bergkamen, Germany, Federal Republic of Henderson, David, Berlin, Germany, Federal Republic of Contain Aktiengesellschaft, Berlin and Bergkamen, Germany, Federal Republic of Common Schering Aktiengesellschaft, Berlin and Bergkamen, Germany, Federal Republic of (non-U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION: US 1995-910148 19951218 (6)

RELATED APPLN. INFO:: Continuation-in-part of Ser. No. US 1994-621308, filed on 15 Jun 1994, now abandoned

NUMBER DATE

PRIORITY INFORMATION: DE 1993-3321826 19830615

DE 1994-3441036 19840404

DE 1994-3446661 19841218

DOCUMENT TYPE: Utility

FILE SEGMENT: Granted

Schenkman, Leonard

ASSISTANT EXAMINER: Lipowsky, Joseph A. Millen & White

NUMBER OF CLAIMS: 41

EXEMPLARY CLAIM: 18

LINE COUNT: 310

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB 13.alpha.-alkylgonanes of formula I #STR1## where R is an acyl radical with as many as 10 C-atoms, and

X is an oxygen atom or the grouping N--OH,

have a strong antigestagenic effect and can be used for postcoital fertility control.

IT 96285-40-49 96285-50-69

(prepn. of, as postcoital contraceptive)

RN 96285-40-4 USPATFULL

CN 19-Norpregna-4, 9-diene-3, 20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-, (11.beta., 13.alpha.)- (9CI) (CA INDEX NAME)

=> d ibib ab hitstr 1-33

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L5 ANSWER 1 OF 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 2002:869589 CAPLUS
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DOCUMENT NUMBER:

2002:869589

137.346927
Implantation rates after in vitro fertilization, and treatment of infertility and early pregnancy loss with a nitric oxide donor or substrate alone or in combination with progesterone, and a method for contraception with nitric oxide inhibitors in combination with antirogestims or other agents Chwalisz, Krzysztof; Garfield, Robert E.

Germany
U.S. Pat. Appl. Publ., 15 pp., Division of U.S. Ser.
No. 162,446.
CODEN: USXXCO
Patent

INVENTOR(S): PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

APPLICATION NO. DATE PATENT NO. KIND DATE

PATENT NO. KIND DATE APPLICATION NO. DATE

US 2002169205 A1 20021114 US 2002-43232 20020114

PRIORITY APPLN. INFO: US 1998-162446 A3 19980299

AB A method is provided for the improvement of implantation rates and/or pregnancy rates in a female mammal, comprising administering to a female mammal in whom pregnancy is desired an effective amt. of (a) a nitric oxide synthase substrate, a nitric oxide donor, or both, optionally in combination with (b) a progestin, and, (c) optionally, in further combination with an estrogen. A method is also provided for fertility control for a female mammal, comprising administering to a female mammal in whom pregnancy is not desired and at risk for becoming pregnant an effective amt. of nitric oxide synthase inhibitor in combination with an antiprogestin. Pharmaceutical compns. are also provided.

IT 126784-99-4, CDB 2914

(Biological study): USES (Uses)

(antiprogestin, method for contradeption with nitric oxide inhibitors in combination with antiprogestins or other agents)

RN 12678-99-4 CABLUS

CN 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4-(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L5 ANSWER 2 OF 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 2002:211446 CAPLUS
DOCUMENT NUMBER: 137:28399
TITLE: CDR-4124

137:28399
CDB-4124 and its putative monodemethylated metabolite, CDB-4453, are potent antiprogestins with reduced antiglucocorticoid activity: in vitro comparison to mifepristone and CDB-2914
Attardi, Barbara J., Burgenson, Janet; Hild, Sheri A.; Reel, Jerry R.; Blye, Richard P.
Molecular Endocrinology Laboratory, BIOQUAL, Inc., Rockville, MD, 20850, USA
Molecular and Cellular Endocrinology (2002), 188(1-2), 111-123
CODEN: MCEND6; ISSN: 0303-7207
Elsevier Science Ireland Ltd.
Journal

AUTHOR (S):

CORPORATE SOURCE:

PUBLISHER:

CODEN: MCENDG; ISSN: 0303-7207

Elsevier Science Ireland Ltd.

CODEN: MCENDG; ISSN: 0303-7207

Elsevier Science Ireland Ltd.

Journal

To obtain selective antiprogestins, we have examd. the in vitro

antiprogestational/antiglucocorticoid properties of two novel compds.,

CDB-4124 and the putative monodemethylated metabolite, CDB-4453, in

transcription and receptor binding assays and compared them to CDB-2914

and mifepristone. All four antiprogestins bound with high affinity to

rabbit uterine progestin receptors (PR) and recombinant human PR-A and

PR-B (rhRP-A, rhPR-B) and were potent inhibitors of R5020-induced

transactivation of the PREZ-tk-luciferase (PREZ-tk-LUC) reporter plasmid

and endogenous alk, phosphatase prodn. In T470-CO human breast cancer

cells. None of these compds. exhibited agonist activity in these cells.

Induction of luciferase activity was potentiated about five-fold by

8-Br-cAMB under basal conditions and to the same extent in the profence of

the PR antagonists. Mifepristone bound to rabbit thymic glucocofticoid

receptors (GR) with approx. twice the avidity of the COB antip/ogestins.

Inhibition of GA-mediated transcription of PREZ-tk-LUC was apsessed in

Hep62 human hepatoblastona cells. Mifepristone exhibited gfeater

antiglucocorticoid activity than CGB-2914, 4124, and 4453 about 12-, 22-,

and 185-fold, resp. Thus, while there was a good correlation between

binding to PR and functional activity of these antiprogestins, GR binding

was not predictive of their glucocorticoid antagonist activity. In

agreement with our in vivo results, COB-4124 and CB-4453, as well as

CDB-2914, are potent antiprogestins in vitro, but show considerably less

antiglucocorticoid activity than mifepristone.

198414-31-2, CDB-4124 and Statis-28-0, CDB 4453

RL: PAC (Pharmacological activity): THU (Therapeutic use); BIOL

(Biological study); USES (Uses)

(CDB-4124 and putative monodemethylated metabolite, CDB-4453, are

potent antiprogestins with reduced antiglucocorticoid activity in

transcription DOCUMENT TYPE: LANGUAGE: AB To obtain

Absolute stereochemistry.

ANSWER 1 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued)

ANSWER 2 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued)

65416-28-0 CAPLUS 19-Norpregna-4,9-diene-3,20-dione, 17-{acetyloxy}-21-methoxy-11-{4-(methylamino)phenyl}-, (11.beta.)- (9CI) (CA INDEX NAME)

126784-99-4, CDB-2914
RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (comparison compd.; CDB-4124 and putative monodemethylated metabolite, CDB-4453, are potent antiprogestins with reduced antiglucocorticoid activity in transcription and receptor binding assays) 126784-99-4 CAPLUS
19-Notpreagna-4, 9-diene-3, 20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 36 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

LS ANSWER 2 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued)

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L5 ANSWER 3 OF 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 2001:747811 CAPLUS
135:304062
TITLE: 315:304062
Preparation of 17. alpha.-substituted-11.beta.-
substituted-4-aryl and 21-substituted
19-norpregna-4, 9-diene-3, 20-diene derivatives as new
antiprogestational agents
Xim, Hyun K. B Blye, Richard P.; Rao, Pemmaraju N.;
Cessac, James W.; Acosta, Carmie K.; Simmons, Anne
Marie

PATENT ASSIGNEE(S): Secretary of Health and Human Services, USA
FOT Int. Appl., 171 pp.
CODEN: PIXXO2

CODEN: PIXXO2

CODEN: PIXXO2

CODEN: PIXXO2

CODEN: PIXXO2

CODEN: PIXXO2

PATENT INFORMATION:

PATENT INFORMATION:

PATENT INFORMATION:

PATENT NO. XIND DATE APPLICATION NO. DATE

WO 2001074840 A2 20011011 WO 2001-US8681 20010316

WO 2001074840 A3 20020502

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GB, GH, GH, HR,
HU, ID, IL, IN, IS, JP, XE, KG, XP, KR, XZ, LC, LK, LR, LS, LT,
LU, LV, AA, MD, MG, MK, MM, HW, MK, MZ, NO, NZ, PL, FT, RO, RU,
SD, SE, SG, SI, SK, SL, TJ, TH, TR, TT, TZ, UA, UG, US, UZ, YM,
RW: GH, GH, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
BJ, CF, CG, CI, CH, GA, GN, GW, ML, MR, NE, SN, TD, TG
AU 2001045849 A5 20011015 AD 2001-25895 A 20010316

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
IE, SI, LT, LY, FI, RO, MK, CY, AL, TR
PRIORITY APPLN. INFO::

WS 2001-258681 W 20010316

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
IE, SI, LT, LY, FI, RO, MK, CY, AL, TR
PRIORITY APPLN. INFO::

WS 2001-258681 W 20010316

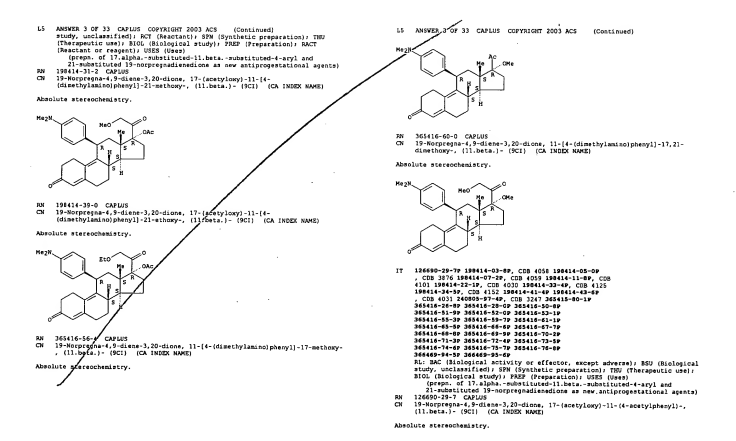
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IE, SI, LT, LY, FI, RO, MK, CY, AL, TR
PRIORITY APPLN. INFO::

WS 2001-258681 W 20010316

R: AT, BE, CH, DE, DK, SE, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
IE, SI, LT, LY, FI, RO, MK, CY, AL, TR
PRIORITY APPLN. INFO::

WS 2001-258681 W 20010316

R: AT, BE, CH, DE, DK, SE, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
IE, SI, LT, LY, FI, RO, MK, CY, AL, TR
P
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L5 ANSWER 3 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued

RN 198414-03-8 CAPLUS
CN 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-21-fluoro-, (11.beta.)- (9C1) (CA INDEX NAME)

Absolute stereochemistry.

RN 198414-05-0 CAPLUS
CN 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-21-chloro-11-[4-ddientylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 198414-07-2 CAPLUS
CN 19-Norpregna-4,9-diene-3,20-dione, 17,21-bis(acetyloxy)-11-[4-(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

L5 ANSWER 3 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued) NAME)

Absolute stereochemistry.

RN 198414-34-5 CAPLUS
CN 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-21-hydroxy-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 198414-41-4 CAPLUS
CN 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-21-methoxy-, 3-oxime, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry unknown.

RN 198414-43-6 CAPLUS

L5 ANSWER 3 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued) Absolute stereochemistry.

RN 198414-11-8 CAPLUS
CN 19-Notpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-21-(acetylthio)-11-[4-(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 198414-22-1 CAPLUS
CN Estra-4,9-dien-3-one, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-17-(1-oxopropyl)-, (11.beta.,17.alpha.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

RN 198414-33-4 CAPLUS
CN 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-21-(3-cyclopentyl-1-oxopropoxy)-11-[4-(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX

L5 ANSWER 3 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued)
CN 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-21-bromo-11-(4-(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 240805-97-4 CAPLUS
CN 19-Norpregna-4,9-diene-3,20-dione, 11-[4-(dimethylamino)phenyl]-17-(1-oxopropoxy)-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 365415-80-1 CAPLUS
N 19-Norpreyna-4,9-diene-3,20-dione, 17,21-bis(acetyloxy)-11-[4-(dimethylamino)phenyl]-, 3-oxime, (11.beta.)- (9C1) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry unknown.

RN 365416-26-8 CAPLUS
CN 19-Norpregna-4,9-diene-3,20-dione, 11-[4-(dimethylamino)phenyl]-17,21-dimethoxy-, 3-oxime, (11.beta.)- (9CI) (CA INDEX NAME)

L5 ANSWER 3 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued)

Absolute stereochemistry. Double bond geometry unknown.

365416-28-0 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-21-methoxy-11-{4-(methylamino)phenyl}-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

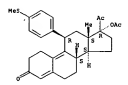
365416-50-8 CAPLUS 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4-acetylphenyl)-21-(acetylthio)-, (11.beta:)- (9CI) (CA INDEX NAME)

365416-51-9 CAPLUS

ANSWER 3 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued) L5

365416-55-3 CAPLUS 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(methylthio)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



365416-59-7 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17,21-bis(acetyloxy)-11-{4-acetylphenyl}-, (11.beta.}- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

365415-61-1 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 11-[4-(dimethylamino)phenyl]-17-ethoxy-21-methoxy-, (11.beta.)- (SCI) (CA INDEX NAME)

Absolute stereochemistry.

365416-65-5 CAPLUS 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4-acetylphenyl)-21-

ANSWER 3 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued) 19-Norpregna-4,-9-diene-3,20-diene, -11-(4-acetylphenyl)-17,21-dimethoxy-, (11.beta.)- (9C1) (CA INDEX NAME)

Absolute stereochemistry.

365416-52-0 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-[2-(dimethylamino)ethoxy]phenyl]-21-methoxy-, (11.beta.)- (9CI) (CA INDEX NAME)

365416-53-1 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-21-methoxy-11-(4-[2-(1-piperidinyl)ethoxy]phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 3 OF 33 CAPLUS COPYRIGHT 2003 ACS methoxy-, (11.beta.)- (9CI) (CA INDEX NAME)

365416-66-6 CAPLUS 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-21-methoxy-11-[4-[2-(1-pyrcolidniy)]ethoxy]phenyl]-, (11.beta.)- (SCI) (CA INDEX NAME)

Absolute stereochemistry.

365416-67-7 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl)-21-(1-oxopropoxy)-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

365416-68-8 CAPLUS 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-

L5 ANSWER 3 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued) (dimethylamino)phenyl]-21-[(methoxyacetyl)oxy]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 365416-69-9 CAPLUS
CN 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-21-[(methoxycarbonyl)oxy]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 365416-70-2 CAPLUS
CN 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-21-(ethenyloxy)-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L5 ANSWER 3 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued) Absolute stereochemistry.

RN 365416-74-6 CAPLUS
CM 19-Norpregna-4,9-diene-3,20-dione, 11-[4-(dimethylamino)phenyl]-17,21-bis(formyloxy)-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 365416-75-7 CAPLUS
CN Glycine, N,N-dimethyl-, (11.beta.)-17-(acetyloxy)-11-[4-(dimethylaminojphenyl)-3,20-dioxo-19-norpregna-4,9-dien-21-yl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 365416-76-8 CAPLUS
CN 19-Norpregna-4,9-diene-3,20-dione, 11-[4-(dimethylamino)phenyl]-17-methoxy-, 3-oxime, (11.beta.)- (9CI) (CA INDEX NAME)

L5 ANSWER 3 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued

RN 365416-71-3 CAPLUS
CN 19-Norpregna-4, 9-diene-3, 20-dione, 11-[4-(dimethylamino)phenyl]-21-(ethenyloxy)-17-methoxy-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

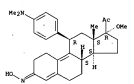
RN 365416-72-4 CAPLUS
CN 19-Norpregna-4,9-diene-3,20-dione, 11-[4-(dimethylamino)phenyl]-21-(ethenyloxy)-17-ethoxy-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 365416-73-5 CAPLUS

Not 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4-idimetylamino)phenyl]-21-thiocyanato-, (11.beta.)- (9CI) (CA INDEX NAME)

L5 ANSWER 3 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued) Absolute stereochemistry.
Double bond geometry unknown.



RN 366469-94-5 CAPLUS
CN 19-Norpregna-4, 9-diene-3, 20-dione, 11-[4-(dimethylamino) phenyl]-17(formyloxy)-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry

RN 366469-95-6 CAPLUS
CN 19-Norpregna-4,9-diene-3,20-dione, 11-[4-(dimethylamino)phenyl]-17-[(1-oxoheptyl)oxy]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 126784-99-4, CDB 2914
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(Uses)
(prepn. of 17.alpha.-substituted-11.beta.-substituted-4-aryl and 21-substituted 19-norpregnadienedione as new antiprogestational agents)

ANSVER 3 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued) 126784-99-4 CAPLUS 19-Nocpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4-(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

365416-20-2P 365416-21-3P

365416-20-29 365416-21-39 Regret Reactant), PREP (Preparation), RACT (Reactant) or reagent) (prepn. of 17.alpha.-substituted-11.beta.-substituted-4-aryl and 21-substituted 19-norpregnadienedione as new antiprogestational agents) 365416-20-2 CAPLUS 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-21-{(chloroacetyl)oxy}-11-(4-(dimethylamino)phenyl}-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

365416-21-3 CAPLUS

Joseffor 21-3 CAPUS
19-Norpregnat4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-21-[(iodoacetyl)oxy]-, (11.beta.)- (9CI) (CA INDEX NAME)

L5 ANSWER 4 OF 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 2001:489415 CAPLUS
DOCUMENT NUMBER: 135:61476
TITLE: Process

135:61476
Process for the preparation of 17.alpha.-acetoxy11.beta.-[4-N,N-(dimethylamino)phenyl]-21-methoxy-19norpregna-4,9-diene-3,20-dione, intermediates useful in the process, and processes for preparing such intermediates

INVENTOR (S):

intermediates

Kim, Hyun Koo Rao, Pemmaraju N., Cessac, James W.;

Simmons, Anne Marie

United States Dept. of Health and Human Services, USA

PCT Int. Appl., 50 pp.

CODEN: PIXXO2

Patent

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE:

Patent

LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

WO 2001047945 A1 20010705 WO 2000-US35479 20001229

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BB, BR, BR, BE, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, K, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, KK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

RW: GH, GH, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, TI, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

AU 2001026048 A5 20010709 AU 2001-26048 20001229

EP 1242444 A1 20020925 EP 2000-989551 20001229

ER 1AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR

US 200306046 A1 20030327 US 2002-169139 20020627

ORITY APPLM. INFO: US 1999-173470F P 19991229

ER SOURCE(S): CASPEACT 135:61476

A process for prepg. the antiprogestational agent, 17. alpha.—acetoxy-11. beta.—[4-N, N-(dimethylamino)phenyl]-21-methoxy-19-norpregna-4, 9-dien -3, 20-dione (I), internediates was described. I was prepd. via a multistep synthetic sequence starting from cynanohydrin II. The synthetic sequence involved replacing the cyanohydrin group of II with a chloroacetyl group and a hydroxyl group; deacetylating the resulting compd.; selectively methylating the Paulting compd. selectively group of the resulting compd. selectively group of the resulting compd. selectively group of the resulting compd. selectively methylating the 20-ktog group of the resulting compd. selectively worldizing the epoxidizing the resulting compd.; selectively oxidizing the epoxidizing the resulting compd. selectively oxidizing the epoxidizing the resulting compd. selectively oxidizing the epoxidizing the resulting compd. selectively oxidizing the selectively indicated the selectively oxidizing the colone, intermediates useful in the process for the prepn. of 17. alpha.—ac PATENT NO. PRIORITY APPLN. INFO.: OTHER SOURCE(S):

L5 ANSWER 3 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued)

365416-27-9P
RL: SPN (Synthetic preparation); PREP (Preparation)
(prepn. of 17.alpha.-substituted-11.beta.-substituted-4-aryl and
21-substituted 19-norpregnadienedione as new antiprogestational agents)
365416-27-9 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17,21-bis(acetyloxy)-11-(4(methylamino)phenyl)-, (11.beta.)- (9CI) (CA INDEX NAME)

ANSWER 4 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued) (dimethylamino)phenyl]-21-methoxy-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT:

THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 5 OF 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 2001:168581 CAPLUS
DOCUMENT NUMBER: 134:361485
TITLE: Effect of a 17.alpha.-(3-Hydroxypropyl)-17.beta.acctyl Substituent Pattern on the Glucocorticoid and
Progestin Receptor Binding of 11.beta.-Arylestra-4,9dien-3-ones
AUTHOR(S): Cook, C. Edgar Raje, Prasady Lee, David Y.-W.,
Kepler, John A.
CORPORATE SOURCE: Chemistry and Life Sciences, Research Triangle
Institute, Research Triangle Park, NC, 27709-2194, USA
Organic Letters (2001), 3(7), 1013-1016
COOEM. ORLEF7, ISSN: 1523-7060
American Chemical Society
DOCUMENT TYPE: Journal
LANGUAGE: English
AB Replacing the 17.alpha.-acetoxy substituent in an antiprogestational
17.beta.-acetyl-11.beta.-arylestra-4,9-dien-3-one by 3-hydroxypropyl
significantly diminished glucocorticoid receptor binding with little
effect on progestin receptor binding.
112678-99-4, RT 3021-012
RL: BPR (Biological process)) BSU (Biological study, unclassified), PRP
(Properties): BIOL (Biological study) PROC (Process)
((hydroxypropyl) acetyl substituent pattern effect on glucocorticoid and
progestin receptor binding of arylestradienones)
RN 126784-99-4 CAPLUS
CN 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4(dimethylaminophenyl]-, (11.beta.)- (9C1) (CA INDEX NAME)

Absolute stereochemistry

REFERENCE COUNT:

THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 6 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued)

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 6 OF 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 2000:880967 CAPLUS
134:33012
TITLE: Pharmaceutical formulations containing hormones for treating postmenopausal and perimenopausal women
Hartin, Kathryn A., Crowley, William F., Jr.
General Hospital Corp., USA
COENT TYPE: COENT PIXXOZ
DOCUMENT TYPE: LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE

WO 200074684 Al 20001214 WO 2000-US40061 20000602

W: As, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, LV, MA, HD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, FT, RO, RU, SD, SE, SG, S1

RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, CF, CG, C1, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

EP 1187618 Al 2020320 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO

JP 2003501390 T2 20030114

PRIORITY APPLIN. INFO: 19990604

WO 2000-USA0061 10000602

RAB Pharmaceuters 1

R: AT. BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO
JP 2003501390 T2 20030114 JP 2001-501220 200006602
SMITY APPLN. INFO.: US 1999-1374409 P 19990604
WO 2000-US40061 W 20000602
Pharmaceutical formulations contg. Various combinations of an estrogen, a progestin, an androgen, a selective estrogen receptor modulator, a selective androgen receptor modulator, and/or a selective progestin receptor modulator for use in treating postmenopausal or perimenopausal vomen are described. The estrogen is selected from the group consisting of, e.g., conjugate destrogens, estratified estrogens, estradiol valerate, estradiol. The androgen is selected from the group consisting of, e.g., progesterone, 17-hydroxyprogesterone, and 19-nortestosterone derivs. The hormones can be administered at 0.01 .mu.g/kg-4 mg/kg (estrogen), 0.01 .mu.g/kg-5 mg/kg (androgen), and 0.0-200 mg/kg (progestogen) via transdermal. buccal, oral, intravaginal, etc., routes. 126784-99-4, CDB2914
RL: THU (Therapeutic Use), BIOL (Biological study); USES (Uses) (pharmaceutical formulations contg. hormones for treating postmenopausal and perimenopausal women) 126784-99-4 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L5 ANSWER 7 OF 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 2000:470069 CAPLUS
COCUMENT NUMBER: 133:208033
TITLE: A practical large-scale synthesis of 17.alpha.-acetoxy-11.beta.-(4-N,N-dimethylaminophenyl)19-norpregna-4,9-diene-3,20-dione (CDB-2914)
Rao, P. N., Acosta, C. K., Bahr, M. L., Burdett, J. E.; Cessac, J. W., Morrison, P. A.; Kim, H. K.
Department of Organic Chemistry, Southwest Foundation for Biomedical Research, San Antonio, TX, 78245-0549, USA
SOURCE: Steroids (2000), 65(7), 395-400
CODEN: STEDAM; ISSN: 0039-128X
PUBLISHER: Elsevier Science Inc.
DOCUMENT TYPE: Journal
LANGUAGE: Anew practical synthesis of 17.alpha.-acetoxy-11.beta.-(4-N,N-dimethylaminophenyl)-19-norpregna-4,9-diene-3,20-dione (CDB-2914) is described. The synthesis gives easily isolable solids at all steps and is amenable to large-scale process.

IT 126784-99-4C, CDB-2914
RL: SFN (Synthetic preparation); PREP (Preparation)
(practical large-scale synthesis of CDB-2914)
RN 126784-99-4 CAPLUS
CN 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylaminophenyl)-1, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT:

THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 8 OF 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 2000:381156 CAPLUS
133:129998
Circulating concentrations of the antiprogestins CDB-2914 and mifepristone in the female rhesus me following various routes of administration Larner, J. M.; Real, J. R.; Blye, R. P.
CORPORATE SOURCE: Bioqual, Inc., Rockville, Mp. 20850, USA
Human Reproduction (2000), 15(5), 1100-1106
CODEN: HUREER; ISSN: 0268-1161
Oxford University Press

CODEN: HURREER ISSN: 2056-1161

ISHER: Oxford University Press

MENT TYPE: Journal

UNGE: English

The overall aim of these studies was to investigate the oral and i.m. bicavailability of CDB-2914 in intact female thesus monkeys, and to compare the serum concns. of CDB-2914 with that of mifepristone following oral administration. In the first study, a SO mg bolus of CDB-2914 per monkey was administered i.v., orally or i.m. The area under the serum concn.-time curve for 72 h (AUCO-72) following i.v. injection was 18 320.+-2718 mg/mb.bul.h, and that for oral administration was 10 464.+-3248 mg/mb.bul.h. Thus, the oral bicavailability of CDB-2914 equiv was 561. In the second study, the serum concns. of CDB-2914 equiv was 621. In the second study, the serum concns. of CDB-2914 equiv was 621. In the second study, the serum concns. of CDB-2914 equiv was 621. In the second study, the serum concns. of CDB-2914 equiv was 621. In the serum concns. (Cmax) and suspending vehicle (ASV), the mean peak serum concn. (Cmax) of CDB-2914 equiv (192.+-64 ng/ml) occurred at 5.+-1 h, whereas the Cmax of mifepristone equiv. (82.+-25 mg/ml) occurred at 3.+-1 h. Following administration in gelatin capsules (35 mg/monkey), the Cmax of CDB-2914 equiv (129.+-24 ng/ml) occurred at 3.+-1 h. The serum concn. (AUCO-120 h) of CDB-2914 equiv va 47.- or 5.3-fold greater than that of mifepristone equiv. When administred orally in ASV or gelatin capsules resp. The serum protein binding characteristics of CDB-2914 ere also studied. CDB-2914 bound to human .alpha.1-acid glycoprotein (AAG), but not with as high an affinity as mifepristone. In contrast, neither CDB-2914 nor mifepristone bound with high affinity to AAG, corticosteroid binding globulin in monkey serum. Collectively, these results indicated that CDB-2914 was more efficiently absorbed than mifepristone following oral administration to female rhesus monkeys. PUBLISHER: DOCUMENT TYPE: LANGUAGE: AB The overal

RL: BOC (Biological occurrence): BFR (Biological process): BSU (Biological study, unclassified): BIOL (Biological study): OCCU (Occurrence): PROC (Process)

(Process)
(circulating concns. of antiprogestins CDB-2914 and mifepristone in female rhesus monkey following various routes of administration in relation to binding by serum proteins)
126784-99-4 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L5 ANSWER 9 OF 33
ACCESSION NUMBER:
DOCUMENT NUMBER:
133:129997
A single mid-follicular dose of CDB-2914, a new antiprogestin, inhibits folliculagenesis and endometrial differentiation in normally cycling women
AUTHOR(S):

CORPORATE SOURCE:

CORPORATE SOURCE:

CORPORATE SOURCE:

CORPORATE SOURCE:

STATEMON DO NOT SOURCE SENSE SE

SOURCE: 20892-1583, USA
Human Reproduction (2000), 15(5), 1092-1099
CODEN: HUREER; ISSN: 0268-161

PUBLISHER: Oxford University Press
Journal
LANGUAGE: English
AB Previous studies in women have shown that the antiprogestin mifepristone delays or inhibits folliculogenesis. The purpose of this study was to explore whether a new analog, CDB-2914, has similar effects on folliculogenesis, ovulation, or on subsequent luteal phase endometrial maturation. Forty-four normally cycling, healthy women recorded urine IH and vaginal bleeding during pre-treatment, treatment, and post-treatment cycles. At a lead follicule diam. of 14-16 mm, a single oral dose (10, 50, 100 mg) of CDB-2914 or placebo was given, and daily ultrasound, estradiol and progesterone were obtained until follicular collapse, an endometrial biopsy was obtained S-7 days later. Single doses of CDB-2914 were well tolerated. Mid-follicular CDB-2914 suppressed lead follicle growth, causing a dose-dependent delay in folliculogenesis and suppression of plasma estradiol. At higher doses, a new lead follicle was often recruited. Although luteinized unruptured follicles were obsd. at the 100 mg dose, all women had follicular collapse. There was a significant delay in endometrial maturation after CDB-2914 at 10 doses. The treatment cycle was lengthened by 1-2 wk in 30t at 100, 27% at 50 and 9t at 10 mg. CDB-2914 altered ovarian and endometrial physical, without major effects on menstrual cyclicity and may have therapeutic utility.

11 12678-99-4, CDB-2914

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study), USES (Uses)

(single mid-follicular dose of CDB-2914, new antiprogestin, inhibits folliculogenesis and endometrial differentiation.

(single mid-follicular dose of CDB-2914, new antiprogestin, inhibits folliculogenesis and endometrial differentiation in normally cycling

women)
126784-99-4 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

ANSWER 8 OF 33 CAPLUS COPYRIGHT 2003 ACS

REFERENCE COUNT:

THERE ARE 28 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 9 OF 33 CAPLUS COPYRIGHT 2003 ACS

REFERENCE COUNT:

THERE ARE 30 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

LS ANSWER 10 OF 33
ACCESSION NUMBER:
DOCUMENT NUMBER:
133:53856
CDB-2914: anti-progestational/anti-glucocorticoid profile and post-coital anti-fertility activity in rats and rabbits
Hild, Sheri Ann; Reel, Jerry R.; Hoffman, Loren H.; Blye, Richard P.
BIOQUAL Inc., Rockville, MD, 20850, USA
Human Reproduction (2000), 15(4), 822-829
COOLEN: HUREER; ISSN: 0269-1161
Oxford University Press
Journal

Human Reproduction (2000), 15(4), 822-829
CODEN: HUMBER; ISSN: 0269-1161
USHER: Oxford University Press
UMENT TYPE: Journal
GUAGE: English
Our goal was to det. the endocrine and post-coital anti-fertility activity
of CDB-2914. Concurrent administration of progesterone to rats on day 4
post-mating blocked the anti-fertility activity of a single oral 2 mg dose
of CDB-2914. CDB-2914 did not exhibit progestational activity in the
estradiol-primed immature female rabbit at doses that exhibited
anti-progestational activity. CDB-2914 antagonized exogenous and
endogenous progesterone-stimulated uterine haptoglobin synthesis and
secretion in immature and adult mated rabbits resp. Neither CDB-2914 nor
misperistone exhibited glucocorticoid activity as detd. by thymus
involution in rats; mispristone was twice as potent as CDB-2914 in
antagonizing glucocorticiod action. Post-coital CDB-2914 treatment
resulted in a dose-dependent redn. In implantation sites and pregnancy
rates in rabbits. CDB-2914-induced inhibition of uterine wt. increase,
endometrial glandular arborization and uterine haptoglobin
synthesis/secretion correlated with inhibition of pregnancy in mated
rabbits. A single oral dose of 64 mg CDB-2914/rabbit was effective at
blocking pregnancy when administered on day 4, 5, or 6 post-mating,
whereas 32 mg/rabbit was only partially effective in this regard. These
data demonstrate that CDB-2914 is a potent, orally active anti-progestin
with weak anti-glucocorticoid activity. CDB-2914 inhibited implantation
in adult rats and rabbits demonstrating its potential as a post-coital
contraceptive drug.

(CDB-2914 as antiprogestin with postcoital antifertility activity and
weak anti-glucocorticoid profile in rats and rabbits)

(CDB-2914 as antiprogestin with postcoital antifertility
activity and
weak antiglucocorticoid profile in rats and rabbits)

(DB-2914 as antiprogestin with postcoital antifertility
activity and
weak antiglucocorticoid profile in rats and rabbits)

Absolute stereochemistry.

L5 ANSWER 11 OF 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 1999:576939 CAPLUS
131:199885 CAPLUS
13

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

R: AT, BE, CH, DE, DX, ES, FR, GB, GR, IT, LI, LU, ML, SE, MC, PT,

1E, SI, LT, LV, FI, RO

RR 9908598 A 20011002 BR 1999-8598 19990305

PRIORITY APPLM. INFO:

US 1998-3594 A 19890305

PRIORITY APPLM. INFO:

US 1998-3594 A 19890305

OTHER SOURCE(S):

MARPAT 131:19988

AB 20-Keto-11.beta.-arylsteroids of foremla I (X = 0, (substituted) NOH, H2, OH, etc., R1 = dialkylamino, imidazolyl, pyrolyl, piperidino, etc., R2 = H, halor, R3 = H, Me, halor M4 = H, aryloxy, (substituted) OH, alkyl, etc., R5 = H, alkyl, halo, aryloxy, etc.) are prepd. which exhibit potent antiprogestational activity. Thus, II was prepd. from 17.alpha.-hydroxymethyl-3-methoxy-19-norpregna-1,3,5(10)-trien-20-one and 4-bromo-N.-dimethylaniline in several steps. The affinity of II for the progesterone hormone receptor was IGSO of 0.7 nM.

IT 240803-97-49 240805-98-59 240805-99-69

240806-03-59 240806-03-69

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SNN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (prepn. of 20-keto-11.beta.-arylsteroids with antiprogestational activity)

RN 240805-97-4 (CAPLUS

N 19-Norpregna-4, 9-diene-3, 20-dione, 11-[4-(dimethylamino)phenyl]-17-(1-oxopropoxy)-, (II.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 10 OF 33 CAPLUS COPYRIGHT 2003 ACS

REFERENCE COUNT:

THERE ARE 49 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 11 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued)

240805-98-5 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 11-[4-(dimethylamino)phenyl]-17- .
[(phenylacetyl)oxy]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

240805-99-6 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17-(benzoyloxy)-11-[4-(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry

19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4-methoxyphenyl)-, (11.beta.)- (9CI) (CA INDEX NAME)

ANSWER 11 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued)

240806-06-8 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)-3-fluorophenyl]-; (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 12 OF 33 CAPLUS COPYRIGHT 2003 ACS

244206-53-9 CAPLUS Acetamide, N-[4-([1].beta.)-17-(acetyloxy)-3,20-dioxo-19-norpregna-4,9-dien-11-yl)henyl]-2,2,2-trifluoro-(SCI) (CA INDEX NAME)

Absolute stereochemistry.

244206-49-3P 244206-50-6P 244206-56-2P
RL: SPN (Synthetic preparation); PREP (Preparation)
(synthesis of N-desseth) derivs. of CDB-2914 and mifepristone as substrates for synthesis of radioligands; 244206-49-3 CAPUS
19-Norprepara-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4-aminophenyl)-,(11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

244206-50-6 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(methylmethyl-t3-amino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

ANSWER 12 OF 33 CAPLUS COPYRIGHT 2003 ACS
25SION NUMBER: 1999:416361 CAPLUS
MENT NUMBER: 131:243453

E: Synthesis of N-desmethyl derivatives of
17. alpha. -acetoxy-11.beta. - (4-N, N-dimethylaminophenyl)19-norpregna-4,9-diene-3,20-dione and mifepristone:
substrates for the synthesis of radioligands
Rao, Pemmaraju N. Acosta, C. Kirk: Cessac, James V. Bahr, Martin L., Kim, Byun K.

ORATE SOURCE: Department of Organic Chemistry, Southwest Foundation
for Biomedical Research, San Antonio, TX, 78245-0549,
USA

ICE: Steroids (1999), 64(3), 205-212
CODEN: STEDAM: ISSN: 0039-128X
Elsevier Science Inc.
MENT TYPE: Journal

AUTHOR (S):

CORPORATE SOURCE:

PUBLISHER: DOCUMENT TYPE: LANGUAGE: AB The synthe JISHER: Elsevier Science Inc.

MENT TYPE: Journal

UNGE: English

The syntheses of N-desmethyl derivs. of CDB-2914 and the mono-N-desmethyl

deriv. of mifepristone are described. We also describe the use of the

mono-desmethyl derivs. as substrates for the synthesis of N-tritiomethyl

derivs. of CDB-2914 and mifepristone with high specific activity (ca. 80

Ci/mmoll, which serve as radioligands for RIA.

126784-99-4, CDB-2914

RL: RCT (Reactant): RACT (Reactant or reagent)

(synthesis of N-desmethyl derivs. of CDB-2914 and mifepristone as

substrates for synthesis of radioligands)

126784-99-4 CAPLUS

19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4
(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Absolute stereochemistry.

L5 ANSWER 12 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued) Absolute stereochemistry.

244206-56-2 CAPLUS Acetanide, N-[4-[(11.beta.)-17-(acetyloxy)-3,20-dioxo-19-norpregna-4,9-dien-11-y1)phenyl)-N-methyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT:

THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

LS ANSWER 13 OF 33 CAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER: 1999:154103 CAPLUS

DOCUMENT NUMBER: 130:291788

The novel progesterone receptor antagonists RTI
30:21-012 and RTI 30:21-022 exhibit complex
glucocorticoid receptor antagonist activities:
implications for the development of dissociated
antiprogestins

AUTHOR(S): Vagner, B. L., Pollio, G., Giangrande, P., Webster, J.
C., Breslin, N., Mais, D. E., Cook, C. E., Vedeckis,
V. V., Cidlowski, J. A., McDonnell, D. P.
CORPORATE SOURCE: Endocation of the development of dissociated
university Medical Center, Durham, NC, 27710, USA
Endocrinology (1999), 140(3), 1449-1458

CODEN: ENDOAD, ISSN: 0013-7227

PUBLISHER: Endocrine Society

DOCUMENT TYPE: Journal
LANGUAGE: English

AB The authors have identified two novel compds. (RTI 3021-012 and RTI
3021-022) that demonstrate similar affinities for human progesterone
receptor (PR) and display equiv. antiprogestenic activity. As with most
antiprogestins, such as RU466, RTI 3021-012, and RTI 3021-022 also bind to
the glucocorticoid receptor (GR) with high affinity. Unexpectedly, when
compared with RU466, the RTI antagonists manifest significantly less GR
antagonist activity. This finding indicates that, with respect to
antiglucocorticoid function, receptor binding affinity is not a good
predictor of biol. activity. The authors have detd. that the lack of a
clear correlation between the GR binding affinity is of the RTI compds. and
their antagonist activity reflects the unique manner in which they
modulate GR signaling. Previously, the authors proposed a two step
"active inhibition" model to explain steroid receptor antagonism: (1)
competitive inhibition of agonist binding and (2) competition of the
antagonist bound receptor with that activated by agonists for DNA response
elements within target gene promoters. Accordingly, the authors obed
alone functioned as an active antagonist. RTI 3021-012 and RTI 3021-022,
functioned solely as "competitive antagonist" since they were capable of
high affinity GR binding, but the r

Absolute stereochemistry.

L5 ANSWER 14 OF 33 CAPLUS COPYRIGHT 2003 ACS ACCESSION NUMBER: 1998:646581 CAPLUS DOCUMENT NUMBER: 130:20723

AUTHOR(S): CORPORATE SOURCE: SOURCE:

PUBLISHER

ANSWER 14 OF 33 CAPLUS COPYRIGHT 2003 ACS ESSION NUMBER: 1998:646581 CAPLUS

MENT NUMBER: 130:20723

LE: Antiovulatory and postcoital antifertility activity of the antiprogestin CDB-2914 when administered as single, multiple, or continuous doses to rats single, multiple, or continuous doses to rats single. More and the programment of the antiprogestin CDB-2914 when administered as single. More and the programment of the antiprogestin (1991), 58 (2), 129-136

CODEN: CCPTAY: ISSN: 0010-7824

LISHER: Blowier Science Inc.
MENT TYPE: Journal
SUMGE: Lenglish

The present studies in rats were undertaken to investigate the potential of a new antiprogestin, CDB-2914, for use as an emergency postcoital contraceptive for women. When given orally at noon on the day of proestrus, both CDB-2914 and miferristone displayed dose-dependent antiovulatory activity however, CDB-2914 was about eight times more potent than mifepristone. Both antiprogestins were considerably less potent in blocking ovulation when injected s.c. To evaluate antifertility activity during continuous low dose administration, rats were dosed orally with 0.5 mg of either CDB-2914 or mifepristone daily, commencing on the day of estrus and continuous for 24 days. Females were cohabited with proven fertile males on day 8 of treatment and were removed 1-3 days later after confirmed mating. The pregnancy rate was significantly reduced only in the CDB-2914-treated females; however, the mean no. of normal implantation sites per pregnant rat was significantly reduced only intepristone as compared with the vehicle control group. CDB-2914 was also found to prevent pregnancy when administered orally after mating from days 0.1, 2, or 3 postmating. The formal amiferial promoted and a serious per rate was significantly reduced by mifepristone as compared with the vehicle control group. CDB-2914 was highly effective in blocking pregnancy when given on either day 4 or 5 postmating. Ollectively, these data demonstrate that CDB-2914 is an orally active postcoital antifertilit DOCUMENT TYPE: LANGUAGE: AB The prese

(Uses)
(antiovulatory and postcoital antifertility activity of antiprogestin
(DB-2914 compared to mifepristone as single, multiple, or continuous
doses to rats)
126784-99-4 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

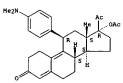
Absolute stereochemistry.

ANSWER 13 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued)

REFERENCE COUNT:

THERE ARE 41 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 14 OF 33 CAPLUS COPYRIGHT 2003 ACS



REFERENCE COUNT:

THERE ARE 36 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

LS ANSWER 15 OF 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 1998:424125 CAPLUS
TITLE: 129:50105
USes of anti-glucocorticoid compounds for the treatment of psychoses or addictive behaviors
Oberlander, Clauder Piazza, Pier Vincenzo
PATENT ASSIGNEE(S): Hoschst Marion Roussel, Fr., Oberlander, Clauder, Piazza, Pier Vincenzo
PCI Int. Appl., 41 pp.
CODEN: PIXXOZ
PATENT INFORMATION: 2

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

RITY APIN. INFO:

FR 1996-15649

ROURCE(S):

MARPAT 129:50105

Glucocorticoid antagonists, except mifepristone, are used as dopamine type
II receptor antagonists to treat psychotic or addictive behavior. Thus,
17.beta.-hydroxy-10.beta.-[(-methylphenyl)methyl]-17.alpha.-[1propynyl]estra-4,9(11)-dien-3-one considerably reduced the response to
morphine in vivo.

126784-99-4

RI: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(use of anti-glucocorticoid compds. as dopamine type II receptor
blocking agents for the treatment of psychoses or addictive behaviors)
126784-99-4

CARUS

19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-{4(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME) OTHER SOURCE(S):

Absolute stereochemistry.

L5 ANSWER 16 OF 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 1998:13308 CAPLUS
DOCUMENT NUMBER: 128:128177
TITLE: 11.beta.-substituted 13.beta.-ethyl gonane derivatives exhibit reversal of antiprogestational activity
AUTHOR(S): Rao, Pemmaraju N., Cessac, James W., Blye, Richard P., Kim, Hyun K.
Department of Organic Chemistry, Southwest Foundation for Biomedical Research, San Antonio, TX, 78245-0549, USA
SOURCE: Steroids (1998), 63(1), 50-57
CODEN: STEDAM; ISSN: 0039-128X
Elsevier Science Inc.
DOCUMENT TYPE: Journal
LANGUAGE: English
AB The syntheses of three 17. alpha.-acetoxy-13.beta'-ethyl-11.beta.-aryl10, 19-dinorpregna-4,9-diene-3, 20 diones from levonorgestrel are described.
Despite their close structural similarity to the antiprogesterone
CDB-2914, one of the compds. exhibits agonistic progestational activity, and the other two compds. are totally inactive.

IT 202062-92-89 202062-93-9P 202062-93-9P 202062-93-9P RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SFN (Synthetic preparation); BIOL (Biological study); PREF (Preparation)
(prepn. of acetoxyethylacyldinorpregnadienediones with reversal of antiprogestational activity)
RN 202062-92-8 CAPLUS

Absolute stereochemistry. Rotation (+).

Absolute stereochemistry. Rotation (+).

202062-93-9 CAPLUS

202002-35-5 CREBUS 18, 19-Dictorpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-13-ethyl-11-[4-(methylthio)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

ANSWER 15 OF 33 CAPLUS COPYRIGHT 2003 ACS

REFERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 16 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued)

202062-94-0 CAPLUS
18,19-Dinorpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4-acetylphenyl)13-ethyl-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

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L5 ANSWER 17 OF 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 1997:745947 CAPLUS
1171LE: 19047
INTELL INTE
                                DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
PATENT NO. KIND DATE APPLICATION NO. DATE

WO 9741866 Al 19971113 WO 1997-EP2371 19970507

W: AL, AM, AT, AL, AZ, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DX, EE, ES, FI, GB, GE, HU, IL, IS, JP, XE, KG, KP, KR, KZ, LX, LR, LS, LT, LU, LY, MD, MG, MK, MN, MY, KN, NO, NZ, PL, PT, RO, RU, SD, SE, SG, S1, SX, T3, TM, TR, TT, UA, UG, UZ, VN, AM, AZ, BY, KO, XZ, MD, RU, TJ, TM

RW: GH, KE, LS, HW, SD, SZ, UG, AT, BE, CH, DE, DX, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CT, CM, GA, GN, HL, MR, NE, SN, TD, TG

US 6040340 A 20000321 US 1996-646518 19960507

AU 19728947 Al 19971126 AU 1997-29947 19970507

EP 906105 Al 1990107 EF 1997-923032 19970507

ER AT, BE, CH, DE, DX, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI

CN 1218402 A 19990602 CN 1997-194452 19970507

BR 9708980 A 19990803 BR 1997-8980 19970507

BG 62953 BI 20001229 BC 1998-102881 19981029

NO 9805204 A 19990106 NO 1998-5204 19981106

KR 2000010833 A 20000225 KR 1998-708974 19981106

PRIORITY APPLIN. INFO: US 1996-66518 A 19960507
                                                                            R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI
CN 1218402 A 19990602 CN 1997-194452 19970507
BR 9708980 A 19990803 BR 1997-8980 19970507
JP 2000510462 T2 20000815 JP 1997-539553 19970507
BG 62953 BI 20001229 BG 1998-102881 19981029
NO 9805204 A 19990106 NO 1998-5204 19981106
KR 2000010833 A 20000025 KR 1998-708974 19981106
KR 1998-10807 WO 1997-EP2371 W 19970507
A method is provided for the improvement of implantation rates and/or pregnancy rates in a female mammal, comprising administering to a female mammal whom pregnancy is desired an effective amt. of: (a) anitric oxide synthase substrate, a nitric oxide donor, or both, optionally in combination with, (b) a progestin, and, (c) optionally, in further combination with, (b) a progestin, and, (c) optionally, in further control for a female mammal; comprising administering to a female mammal in whom pregnancy is not desired and at risk of becoming pregnant an effective amt. of nitric oxide synthase inhibitor in combination with an antiprogestin. Pharmaceutical compns. are also provided.

126784-99-4, CDB2914
RL: RAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(fettility control using a nitric oxide synthase inhibitor in
                                                                                    (Uses)
(fertility control using a nitric oxide synthase inhibitor in combination with an antiprogestin)
126784-99-4 CAPLUS
13-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-
                        L5 ANSWER 18 OF 33 CAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER: 1997:740250 CAPLUS

1097:740250 CAPLUS

11711E: 27:358992

Preparation of 21-substituted progesterone derivatives as new antiprogestational agents

Kim, Hyun K.; Blye, Richard P.; Rao, Pemmaraju N.;

Cessac, James W.; Acosta, Carmie K.

United States Dept. of Health and Human Services, USA;

Xim, Hyun K.; Blye, Richard P.; Rao, Pemmaraju N.;

Cessac, James W.; Acosta, Carmie K.

PCT Int. Appl., 65 pp.

COEN: PIXXO2

Patent
                           CODEN: 1
LANGUAGE: Patent
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
                     APPLICATION NO. DATE
                                                                                    PATENT NO.
                                                                                                                                                                                                                                                                   KIND DATE
                   R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IZ, FI 
AT 194358 E 20000715 AT 1997-923523 19970430 
JP 200050396 T2 20000725 JP 1997-539222 19970430 
US 2002025951 AT 20000725 US 1999-180132 19990430 
US 2002025951 AT 20020228 US 1999-180132 19990530 
PRIORITY APPLIN. INFO: US 1996-18628P P 19960501 
WO 1997-US7373 W 19970430 
OTHER SOURCE(S): HARPAT 127:358992 AB Progesterone derivs. of formula I (R1 = OMe, SMe, NMe2, NHMe, CHO, Ac, CHONCH3; R2 = halo, slkyl, acyl, OM, alkowy, etc., R3 = OH, alkyl, alkoxy, acyloxy; R4 = H, alkyl; X = O, (substituted) NOH) are prepd. as antiprogestational agents. The present invention provides methods wherein the compds. of formula I are advantageously used, inter alia, to antagonize endogenous progesterones to induce menses; to treat endometricalis; to treat dysmenorchea; to treat endocrine hormone-dependent tumors; to treat uterine fibroids; to inhibit uterline endometrial proliferation; to induce labor; and for contraception. Thus, II was prepd. from 3,3-ethylenedioxy-17.beta.-cyano-17.alpha-hydroxyestra-5(10),9(11)-diene and 4-bromon-N,N-dimethylaniline in 9 steps. II showed 2.79 times the antiprogestational potency in the antiClauberg test compared to COB-2914.

IT 198414-07-2P 198414-31-2P RL BAC (Silodgical activity or effector, except adverse); BSU (Biological study); PREP (Preparation); RACT (Reactant) reagents) USES (Uses) (prepn. of progesterone derivs. as antiprogestational agents)
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ANSVER 17 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued) (dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 18 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued) 198414-07-2 CAPLUS 19-Norpregna-4,9-diene-3,20-dione, 17,21-bis(acetyloxy)-11-{d-(dimethylamino)phenyl}-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

198414-31-2 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-21-methoxy-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

198414-03-8F 198414-05-0F 198414-11-8F
198414-32-9F 198414-33-4F 198414-34-5F
198414-39-0F 198414-43-6F
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SFN (Synthetic preparation); THU (Therapeutic use);
BIOL (Biological study); PREF (Preparation); USES (Uses)
(prepn. of progesterone derivs. as antiprogestational agents)
198414-03-8 CAPLUS
198414-03-8 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-{4-(dimethylamino)phenyl]-21-fluoro-, (11.beta.)- (9CI) (CA INDEX NAME)

ANSWER 18 OF 33 CAPLUS COPYRIGHT 2003 ACS

CH2F

198414-05-0 CAPLUS 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-21-chloro-11-{4-(dimethylamino)phenyl}-, {11.beta.}- {9CI} (CA INDEX NAME)

198414-11-8 CAPLUS
19-Worpregna-4, 9-diene-3, 20-diene, 17-(acetyloxy)-21-(acetylthio)-11-[4-(diesthylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

198414-22-1 CAPLUS Estra-4,9-dien-3-one, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-17-(1-oxopropyl)-, (11.beta.,17.alpha.)- (9CI) (CA INDEX NAME)

ANSWER 18 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued)
198414-39-0 CAPLUS
19-Notpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-21-ethoxy-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

198414-43-6 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-21-bromo-11-[4-(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

198414-40-3P 198414-41-4P
RL: SPN (Synthetic preparation); PREP (Preparation)
(prepn. of progesterone derivs. as antiprogestational agents)
198414-40-3 CAPUS
198414-40-3 CAPUS
19-Norpregna-4,9-diene-3,20-dione, 17,21-bis(acetyloxy)-11-[4-(dimethylamino)phenyl]-, 3-oxime, (3E,11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

L5 ANSWER 18 OF 33 CAPLUS COPYRIGHT 2003 ACS Absolute stereochemistry. Rotation (+). (Continued)

198414-33-4 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-21-(3-cyclopentyl-1-oxopropoxy)-11-[4-(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

198414-34-5 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-21-hydroxy-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L5 ANSWER 18 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued)

198414-41-4 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-21-methoxy-, 3-oxime, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry unknown.

L5 ANSWER 19 OF 33 CAPLUS COPYRIGHT 2003 ACS ACCESSION NUMBER: 1996:705614 CAPLUS DOCUMENT NUMBER: 125:329114

125:329114
improved preparation of 17.alpha.-acetoxy-11.beta.-(4-N,N-dimethylaminophenyl)-19-norpregna-4,9-diene-3,20-dione and its intermediates
Kim, Hyun K.; Rao, Pemmaraju Narasinha; Burdett, James
E., Jr.; Acosta, Carmie Kirk
United States Dept. of Health and Human Services, USA
PCT Int. Appl. 40 pp.
CODEN: PIXXO2
Patent
English

INVENTOR(S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| | | ENT : | | | | ND | DATE | | | A | PPLI | CATI | ON NO | ο. | DATE | | | |
|------|------|-------|------|------|-----|-----|------|------|-----|-------|------|------|-------|---------|------|------|-----|-----|
| | | | | | | | | | | - | | | | | | | | |
| | . WO | 9630 | 390 | | A. | ? | 1996 | 1003 | | w | 0 19 | 96-U | S366 | Ω | 1996 | 031R | | |
| | | 9630 | | | | | | | | _ | | | | - | | | | |
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| | | w: | | | | | AZ, | | | | | | | | | | | |
| | | | ES, | FΙ, | GB, | GE, | HU, | IS, | JP, | ΚE, | ΚG, | ΚP, | KR, | ΚZ, | LK, | LR, | LS, | LT, |
| | | | LU, | LV, | MD, | MG, | MK, | MN, | MW, | MX. | NO. | NZ, | PL. | PT. | RO. | RU. | SD. | SE. |
| | | | SG. | | | | | | | | | | | | | | | |
| | | RW: | | | MW, | SD, | SZ, | UG, | AT, | BE, | CH, | DE, | DK, | ES, | FI, | FR. | GB. | GR. |
| | | | IE. | IT. | LU. | MC. | NL, | PT. | SE. | BF. | BJ. | CF. | CG. | CI. | CM. | GA. | GN. | MT. |
| | IIS | 5929 | | | | | 1999 | | | | | | | | | | ٠, | |
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| | | 2216 | | | | | | | | | | | | | | | | |
| | | 9653 | | | | | | | | A | U 19 | 96-5 | 3145 | | 1996 | 0318 | | |
| | ΑU | 7168 | 94 | | В: | 2 | 2000 | 0309 | | | | | | | | | | |
| | EP | 8177 | 93 | | A: | 2 | 1998 | 0114 | | Е | P 19 | 96-9 | 0974 | 9 | 1996 | 0318 | | • |
| | | | | | | | DK, | | | | | | | | | | MC. | PT. |
| | | | IE. | | | | | | | | | | , | | , | | , | , |
| DDTA | DITT | APP | | | | | | | | 110 1 | ODE | 4127 | | | 1995 | 0220 | | |
| FRIO | A11. | , Art | mi4. | THEO | • • | | | | | | | | | | 1006 | | | |
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R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI
PRIORITY APPLN. INFO.:

US 1995-413755 A 19950330

OTHER SOURCE(S):

CASREACT 125:329114; MARPAT 125:329114

M

Absolute stereochemistry.

L5 ANSWER 20 OF 33 CAPLUS COPYRIGHT 2003 ACS ACCESSION NUMBER: 1996:540408 CAPLUS DOCUMENT NUMBER: 125:238850

TITLE: AUTHOR(S):

125:23850

Effects of two antiprogestins on early pregnancy in the long-tailed macaque (Macaca fascicularis)

Tarantal, Alice F.: Hendrickx, Andrew G.: Matlin, Stephen A.: Laaley, Bill L.: Gu, Quin-Quin: Thomas, Charles A.A.: Vince, Pamela M.: Van Look, Paul F.A. California Regional Primate Research Center, University of California, Davis, CA, 95616, USA Contraception (1996), 54(2), 107-115

CODEN: CCPTAY: ISSN: 0010-7824

Elsevier

CORPORATE SOURCE:

SOURCE:

PUBLISHER: DOCUMENT TYPE: LANGUAGE:

COLTRIGORY, ISSN: 0010-7824

LISHER: Elsevier

CODEN: CCTTAY, ISSN: 0010-7824

LISHER: Elsevier

MEMT TYPE: Journal

The abortifacient effects of mifepristone and HRP 2000 were compared in gravid long-tailed macaques. Thirty-six animals were studied with treatment administered either by the oral (0.5 or 5.0 mg/kg; N = 5 per antiprogestin per dose) or i.m. (IM) routes (0.5 mg/kg; N = 5 per antiprogestin) on gestational days (GD) 23-26; six vehicle controls were included. Blood samples were collected for assay of progesterone (P4) and each of the antiprogestins (pre-treatment, daily GD 23-28, every other day GD 30-40), and animals were monitored sonos, throughout gestation.

Results of these studies indicated high rates of abortion with IM administration (3/5 mifepristone, 4/5 HRP 2000) and 5.0 mg/kg oral route (4/5, 2/5, resp.), with less effects noted at oral doses of 0.5 mg/kg (2/5, 0/5, resp.). No early abortions were obsd. in the control groups. Following daily IM treatment, peak levels of 8-16 ng/mL mifepristone were detected whereas 6-10 ng/mL of HRP 2000 were noted (DD 26-27). No secum elevels of mifepristone were detected following either of the oral doses whereas serum levels of 2-6 ng/mL HRP 2000 were noted with high dose oral administration. Results of these studies suggest: (1) both antiprogestins are roughly comparable in terminating early pregnancy although HRP 2000 may be more effective when administered IM whereas mifepristone may be more effective wend administered orally. (2) similar levels of biol. activity are seen with the IM and high dose oral dosing regimens, with little or no activity with the oral low dose, and (3) infants resulting from surviving pregnancies were not affected by early gestation exposure. 126784-894

RL: BRP (Biological process) BSU (Biological study, unclassified), TRU (Therapeutic use), BIOL (Biological study), PROC (Process): USES (Usen)

126784-99-4
RL: BPR (Biological process); BSU (Biological study, unclassified); THU
(Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)
(abortifacient effects of antiprogestins in early pregnancy in
long-tailed macaque in relation to dose and administration route)
126784-99-4 CARUS
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

ANSWER 19 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued)

ANSWER 20 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued)

L5 ANSWER 21 OF 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER:
DOCUMENT NUMBER:
1996:498851 CAPLUS
125:238820
16.alpha.-substituted analogs of the antiprogestin RU486 induce a unique conformation in the human progesterone receptor resulting in mixed agonist activity

Vagner, Brandee L., Pollio, Giusepper, Leonhardt, Susani Vani, Mansukh C., Lee, David Y.-V., Imhof, Markus O., Edwards, Dean P., Cook, C. Edgar; McDonnell, Donald P.

CORPORATE SOURCE: Department Pharmacology Molecular Cancer Biology, Duke University Medical Center, Durham, NC, 27710, USA

SOURCE: Proceedings of the National Academy of Sciences of the United States of America (1996), 93(16), 8739-8744

CODEN: PRASAG, ISSN: 0027-8424

PUBLISHER: National Academy of Sciences
Journal Complexes: Thus, alink was established between the structure of a ligand-receptor complex and biol. activity. In this study, the authors have utilized a series of in vitro assays with which to study hPR pharmacol. and have identified a third class of hPR ligands that induce a receptor conformation which is distinct from that induced by agonists or antagonists. Importantly, when assayed on PR-responsive target genes these compda. were shown to exhibit partial agonist activity an activity for estrogen receptor, the overall structure of the ligand. It appears, therefore, that the obsd. differences in the activity of some PR and estrogen receptor; the overall structure of the ligand-receptor complex is influenced by tell context. Thus, as has been shown previously for estrogen receptor, the overall structure of the ligand. It appears, therefore, that the obsd. differences in the activity of some PR and estrogen receptor; the overall structure of the ligand receptor complex is influenced by tell material adding domains of steroid hormone receptors resulting in different biologies.

IT 126784-99-4, RTI 3021-012

RL: BRC (Biological activity or effector, except adverse); BPR (Biological Indice); PRP (Properties); BIOL (Biological Study); PROC (Process)

(16.alpha--substituted an

Absolute stereochemistry.

L5 ANSWER 22 OF 33 CAPLUS COPYRIGHT 2003 ACS ACCESSION NUMBER: 1995:985962 CAPLUS COPYRIGHT 2003 ACS 124:22540 Pharmacol Pharm

124:22540
Pharmaceutical compositions of antiglucocorticoid compounds for treating or preventing symptoms of spontaneous or narcotic-induced withdrawal. Petit. Francis; Philibert, Daniel; Ulmann, Andre Roussel-UCLAF, Fr.
Eur. Pat. Appl., 30 pp.
CODEN: EPXXOW
Patent

INVENTOR(S): PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| P. | ATE | ENT : | NO. | | KIN | D | DATE | | | AP | PLIC | ATI | ON N | ю. | DATE | | | |
|------|-----|-------|------|------|-----|-----|------|------|-----|-------|------|-----|------|-----|------|------|-----|----|
| - | | | | | | - | | | | | | | | | | | * | |
| Ε | ₽ € | 5762 | 03 | | A1 | | 1995 | 1011 | | EP | 199 | 5-4 | 007€ | 54 | 1995 | 0406 | | |
| | | | | BE, | CH, | DE, | DK, | ES, | FR, | GB, | GR, | IE, | IT, | LI, | LU, | NL, | PT, | SE |
| F. | R 2 | 2718 | 354 | | A1 | | 1995 | 1013 | | FR | 199 | 4-4 | 156 | | 1994 | 0408 | | |
| F. | R 2 | 2718 | 354 | | В1 | | 1996 | 0503 | | | | | | | | | | |
| Z. | A S | 9502 | 058 | | A | | 1996 | 0313 | | ZA | 199 | 5-2 | 058 | | 1995 | 0313 | | |
| C. | A 2 | 2146 | 600 | | AA | | 1995 | 1009 | | CA | 199 | 5-2 | 1466 | 00 | 1995 | 0407 | | |
| F. | 1 9 | 9501 | 683 | | A | | 1995 | 1009 | | FI | 199 | 5-1 | 683 | | 1995 | 0407 | | |
| | | 516 | | | A1 | | 1995 | 1019 | | AU | 199 | 5-1 | 6326 | 5 | 1995 | 0407 | | |
| | | | 8017 | | A2 | • | 1995 | 1024 | | JP | 199 | 5-1 | 0707 | 1 | 1995 | 0407 | | |
| н | U 7 | 7146 | θ | | A2 | ! | 1995 | 1128 | | HU | 199 | 5-1 | 019 | | 1995 | 0407 | | |
| C | N 1 | 116 | 929 | | Α | | 1996 | 0221 | | CN | 199 | 5-1 | 0401 | 5 | 1995 | 0407 | | |
| ORT! | TV | ADD | TM ' | INFO | | | | | | TD 10 | 04 4 | 156 | | | 1004 | 0400 | | |

CN 116929 1 19960221 CN 1995-104015 19950407
RRTY APPLM. INFO:
FR 1994-4156 19940408
RR SOURCE(S): MARPAT 124:22540
Antiqlucocorticoid steroids such as mifepristone, onapristone,
lilopristone and related steroids are proposed for the prevention or
treatment of withdrawal syndromes, either spontaneous or pptd. by
narcotics or mixts. of narcotics. These antiqlucocorticoids would be
useful in the withdrawal from morphinomimetics such as heroin, morphine or
methadone as well as cocaine. Pharmacol. activity was demonstrated by the
effect of the antiqlucocorticoids on the stereotypic behavior of mice in
teaponse to narcotics. Spontaneous withdrawal syndrome was induced by
administration of the opicid antagonist, naloxone. An antiprogesterone
activity of the steroids in their action mechanism was eliminated.
Results confirmed the involvement of endogenous glucocorticoids in
morphine withdrawal since this is inhibited by antiglucocorticoids or
addenalectomy. PRIORITY APPLN. INFO.: OTHER SOURCE(S): AB Antiglucocorticoic

adrenalectory.

126784-99-4
RL: THU (Therapeutic use): BIOL (Biological study): USES (Uses)
(RU 486 related: antiglucocorticoid steroids for treatment or prevention of spontaneous opioid or narcotic-induced drug withdrawal

syndrome.)
126784-99-4 CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-{4-(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 21 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued)

ANSWER 22 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued)

ACCESSION NUMBER:

DOCUMENT NUMBER:

AUTHOR(S): CORPORATE SOURCE:

SOURCE:

DOCUMENT TYPE:

ANSWER 23 OF 33 CAPLUS COPYRIGHT 2003 ACS
ESSION NUMBER: 1995:499191 CAPLUS
UNENT NUMBER: 122:256542
LE: The anti-progestin CDB 2914 has no antifertility
effect in male rats
HOR(S): Wang, Christina; Sinha-Hikim, Amiya; Leung, Andrew
Department of Medicine, Cedars-Sinai Medical Center,
Los Angeles, CA, USA
COE: Contraception (1995), 51(3), 215-18
CODEN: CCPTAY; ISSN: 0010-7824
UNENT TYPE: Journal
GUAGE: English
This study examines the effect of an anti-progestin (CDB 2914) with
anti-progestational potencies similar to RU 486 on spermatogenesis, sperm
maturation, and fertility in male rats. Adult male rats of proven
fertility were administered the anti-progestin (10 mg/kg/day) or vehicle
(control group) for 14, 35, and 70 days to study the possible effect of
this compd. on epididymal sperm maturation, post-meiotic sperm
development, spermatogenesis, and fertility, resp. Fertility rates of the
rats were detd. by mating studies. The anti-progestin, CDB 2914, had no
effect on testis or accessory organ vts. epididymal sperm content or
motility, testicular sperm count, spermatogenesis, and fertility of male
rats. This study suggests that anti-progestins, when administered ewe at
higher doses than those used in humans, have no contraceptive effect in
adult male rats.
126784-99-4, CDB 2914
(RI: BAC (Biological activity or effector, except adverse); BSU (Biological
study, unclassified); BIOL (Biological study)
(anti-progestin CDB 2914 has no antifertility effect in male rats)
126784-99-4 CAPIUS
19-Notpregma-4,9-diene-3,20-diene,17-(acetyloxy)-11-[4(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 24 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued)

L5 ANSWER 24 OF 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 1995:86211 CAPLUS
DOCUMENT NUMBER: 122:31745

TITLE: Oxidative demethylation of 4-substituted
N,N-dimethylanilines with iodine and calcium oxide in the presence of methanol
ACOSTA, Kirk Cessac, James W., Rao, P. Narasimha;
Kim, Kyun K.

CORPORATE SOURCE: Dep. Org. Chem., Southwest Foundation Biomed. Res.,
SOURCE: Journal of the Chemical Society, Chemical
Communications (1994), (17), 1995-6
CODEN: JOURNAL OF COMMUNICATION OF COMMUNIC

Absolute stereochemistry.

159681-66-OP
RL: SPN (Synthetic preparation), PREP (Preparation)
(oxidative demethylation of 4-substituted N,N-dimethylanilines with iodine and calcium oxide in methanol)
159681-66-O CAPLUS
19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(methylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L5 ANSWER 25 OF 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 1994:290311 CAPLUS
120:290311 APRUS
TITLE: A comparison of the pregnancy-terminating potencies of three anti-progestins in guinea pigs, and the effects of sulprostone
AUTHOR(S): Poyser, N. L.; Forcelledo, M. L.
CORPORATE SOURCE: Med. Sch., Univ. Edinburgh, Edinburgh, EH8 9JZ, UK
Prostaglandins, Leukotrienes and Essential Fatty Acids
(1994), 50(5), 245-7
CODEN: PLEABU; ISSN: 0952-3278
DOCUMENT TYPE: Journal
LANGUAGE: Brilish
AB The anti-progestins mifepristone, lilopristone (ZX 98734) and HRP 2000
were equipotent at terminating the pregnancy of guinea-pigs during
mid-gestation, although mifepristone was more effective at low doses.
Sulprostone administration on the day following anti-progestin tended to increase the effectiveness of mifepristone and HRP 2000, without
affecting the time interval between the start of the anti-progestin treatment and the day of abortion. It is concluded that, of the three
afferent anti-progestins used, none is more potent than the other two at
terminating pregnancy in the animal model used. The co-administration of
a PGEZ analog tends to increase the effectiveness of the anti-progestin.

1 126784-99-4 CAPLUS
CN 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4(dimethylamino)phenyl)-, (11.beta.)- (SCI) (CA INDEX NAME)

L5 ANSWER 26 OF 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 1993:73787 CAPLUS
DOCUMENT NUMBER: 118:73787
IIILE: Reversal of activity profile in analogs of the antiproquestin RU 486: effect of a 16.alpha.-substituent on progestational (agonist) activity

AUTHOR(S): Cook, C. Edgar, Wani, Mansukh C., Lee, Yue Wei, Fail, Patricia A., Petrow, Vladimir

CORPORATE SOURCE: Research Triangle Inst., Research Triangle Park, NC, 27709-2194, USA
SOURCE: Life Sciences (1993), 52(2), 155-62
CODEN: LIFSAK, ISSN: 0024-3205
DOCUMENT TYPE: Journal LANGUAGE: English
AB RU 486 analogs (1, R = H, OAC; R1 = H, Et, R2 = H, Me) were tested for binding to progestogen receptors and for progestational and antiprogestational activity. The 17-beta-acetoxy analogs showed antiprogestational activity, whereas the 16.alpha.-Et analogs were progestogenic. The analog (R = R = R2 = H) exhibited mixed activity. Exam. of structure-activity relationships in combination with computer aided mol. modeling suggests that a binding interaction of the 16.alpha.-Et group with the progesterone receptor (PR) or the PR-progestin response element complex may play the major role in this reversal of activity profile.

112690-26-4 (28784-99-4
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified), BIOL (Biological study)
(antiprogestogenic activity of, mol. structure in relation to)
NN 126690-26-4 (ACPIUS
CN 19-Morpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4-(dimethylamino)phenyl]-6-methyl-, (6.alpha.,11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

126784-99-4 CAPLUS 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L5 ANSWER 27 OF 33 CAPLUS COPYRIGHT 2003 ACS ACCESSION NUMBER: 1990:198892 CAPLUS DOCUMENT NUMBER: 112:198892 TITLE: Preparation of 11.beta.

112:198892
Preparation of 11.beta.-aryl-19-norsteroids as antiglucocorticoids, progestogens, and antiprogestogens
Cook, C. Edgar: Wani, Mansukh C.; Lee, Yue Wei; Reel, Jerry R.; Rector, Douglas
Research Triangle Institute, USA
PCT Int. Appl., 50 pp.
CODEN: PIXXD2
Patent

INVENTOR(S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE:

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| | PAT | ENT | NO. | | KI | ND | DATE | | | AF | PLI | CATI | ON 1 | NO. | DATE | |
|------|------|------|------|------|-----|-----|------|------|-----|---------------|-------|---------|--------------|-----|------|------|
| | | | | | | | | | | | | | | | | |
| | WO | 8912 | 448 | | A | 1 | 1989 | 1228 | | WC | 19: | 89-U | 527 (| 06 | 1989 | 0623 |
| | | V: | AU, | DK, | JP, | KR, | NO | | | | | | | | | |
| | | RW: | AT, | BE, | CH, | DE, | FR, | GB, | IT, | LU, | NL, | SE | | | | |
| | US | 4954 | 490 | | A | | 1990 | 0904 | | ÜS | 19: | 88-2 | 1050 | 33 | 1988 | 0623 |
| | | | | | | | 1997 | | | | | | | | 1989 | |
| | | | | | | | 1990 | | | | | | | 6 | 1989 | |
| | | | | | | | 1993 | | | | | 09-3 | 0300 | • | 1303 | 0023 |
| | | | | | | | | | | | | | | | | |
| | | | | | | | 1991 | | | E | 19 | 9-9 | 0/94 | 4 | 1989 | 0623 |
| | EP | | | | | | 1997 | | | | | | | | | |
| | | R: | AT, | BE, | CH, | DE, | FR, | GB, | IT, | LI, | LU, | NL, | SE | | | |
| | JP | 0350 | 5582 | | T. | 2 | 1991 | 1205 | | JF | 19 | 9-5 | 0739 | 92 | 1989 | 0623 |
| | JP | 2953 | 725 | | В: | 2 | 1999 | 0927 | | | | | | | | |
| | AT | 1498 | 39 | | E | | 1997 | 0315 | | A1 | 19 | 9-9 | 0792 | 24 | 1989 | 0623 |
| | US | 5073 | 548 | | | | | | | US | | | | | 1990 | |
| | | 9005 | | | | | 1990 | | | | | 90-5 | | | 1990 | |
| | | 1782 | | | | | 1995 | | | | . 13. | , o - 5 | 340 | | 1330 | 1221 |
| | | | | | | | | | | | | | | | | |
| | | 1782 | | | c | | 1996 | | | | | | | | | |
| | | 9003 | | | A | | 1990 | 1221 | | | | 90-3 | | | 1990 | |
| PRIC | RITY | APP | LN. | INFO | . : | | | | ι | JS 19 | 88- | 2105 | 03 | | 1988 | 0623 |
| | | | | | | | | | · · | I O 10 | 20-1 | 1527 | ne. | | 1000 | 0623 |

RRTTY APPLM. INFO::

US 1988-210503

US 1998-210503

IP980623

RR SOURCE(S):

MARPAT 112:198892

The title compds. [Ir Rl = H, alkyl, alkenyl, etc.; R2 = H, R3 = H, alkyl, alkenyl, alkynyl; R4 = H, Mer, F, Clr R6 - H, MeZN, MeO, MeCO, MeS, etc.; X = O, MeON; or R1R2 = bond; or R1R3 - CH2, N:MCH2; or R2R3 = CH2] were preped. Grignard reaction of S.alpha., 6-alpha. = penye-6. alpha. = methyl-3,3:20,20-bis(ethylenedioxy)-19-norpregn-9(11)-en-17.alpha.-ol (prepn. given) with p-Me2NCGH4MgBr followed by 17-O-acetylation and deketalization gave I [R1 = AcO, R2 = R3 = H, R4 = Me, R6 = MeZN, X = O]. The binding affinity of I for progesterone receptor in cytosol obtained from estrogen-primed immature rabbit uterus was 8-80% that of progesterone. Several I had glucocorticoid receptor binding affinities up to 2.5-fold that of dexamethasone, and one compd. had in vivo antiprogestational activity comparable to that of RU-486.

126509-26-49 126509-29-79 126784-99-49

RL: SPN (Synthetic preparation) PREP (Preparation) (prepn. of, as antiglucocorticoid and/or (anti)progestogen) 126509-26-4 CAPLUS

19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4-(dimethylamino)phenyl)-6-methyl-, (6.alpha.,11.beta.)- (9CI) (CA INDEX NAME) OTHER SOURCE(S):

Absolute stereochemistry.

ANSWER 26 OF 33 CAPLUS COPYRIGHT 2003 ACS

ANSWER 27 OF 33 CAPLUS COPYRIGHT 2003 ACS

126690-29-7 CAPLUS 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4-acetylphenyl)-, (11.beta)- (9C1) (CA INDEX NAME)

Absolute stereochemistry.

126784-99-4 CAPLUS 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-, (11.beta.)- (9CI) (CA INDEX NAME)

LS ANSWER 28 OF 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 1989:213172 CAPLUS
DOCUMENT NUMBER: 110:213172
ITILE: 110:213172
INVENTOR(S): 13/(Alpha) - alkylgonanes, their production, and pharmaceutical preparations containing same Neef, Guenter, Wechert, Rudolf; Beier, Sybiller Elger, Walter, Henderson, David
Schering A.-G., Fed. Rep. Ger.
U.S., 5 pp. Cont. of U.S. Ser. No. 621,308.
COODEN: USXXAM
DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 4 FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE

US 4780461 A 19881025 US 1985-810148 19851218

DE 3321826 A1 19841220 DE 1983-3321826 19830615

DE 3413036 A1 19851017 DE 1984-3413036 19840104

DE 3446661 A1 19860619 DE 1983-3321826 19830615

DE 1984-3413036 19840615

DE 1984-3446661 19841218

OTHER SOURCE(S): CASREACT 110:213172/ MARPAT 110:213172

AB 13.alpha.-Alkylgonanes [1: R = Cl-4 acyl, X = 0, NOH; II; R1 = amino; R2 = H, Me, Et; R3 = (substituted) alkyl; R4 = OH, alkowy, alkanoyloxy; or R3R4

- Q: R5 = H, alkyl; III; Z = CH2CH2, CH2CMe2CH2], having antigestagenic activity and useful as postcoital contraceptives, or for triggering abortion and menstruation (no data), are prepd. via photochem.

epimerization of the 13.beta.-gonanes IV. 11.beta.-(4
Dimethylaminomethyl)-17.alpha.-hydroxyr-13.alpha.
methyl-17.beta.-(4-dimethyl-17a.blan.-methyl-17.beta.-(3
hydroxypropyl)-4,9-gonadien-3-one (V) was acetylated with Ac20 in pyridine to give 11.beta.-(4-dimethyl-17a.blan.-methyl-17.beta.-(3
cetoxypropyl)-4,9-gonadien-3-one. A tablet was formulated conts, V 10.0, lactose 140.0, corn starch 69.5, polyvinylpyrrolidone 25 2.5, Aerosil 2.0, and Mg stearate 0.5 mg.

IN 96285-40-4 CR2828-50-6P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of, as postcoital contraceptive)

RN 96285-40-4 CR2IUS

CN 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4
(dimethylaminophenyl]-7, (11.beta.,13.alpha.)- (9CI) (CA INDEX NAME) PATENT NO. KIND DATE APPLICATION NO. DATE

Absolute stereochemistry.

L5 ANSWER 29 OF 33 CAPLUS COPYRIGHT 2003 ACS ACCESSION NUMBER: 1988:529463 CAPLUS DOCUMENT NUMBER: 109:129463 109:129463
New 11-(alkynylphenyl)-substituted 19-nor and
19-nor-D-homo steroids, their formation and
pharmacological activity, and processes for their
preparation
Teutsch, Jean Georges; Klich, Michel; Philibert,
Paniel TITLE:

Teutsch, vean Carlo Daniel Roussel-UCLAF, Fr. Eur. Pat. Appl., 88 pp. CODEN: EPXXDW PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

INVENTOR(S):

| PATENT NO. | KIND | DATE | | APPLICATION NO. | DATE |
|--------------------|---------|-----------|----------|-----------------|----------|
| | | | | | |
| EP 245170 | A1 | 19871111 | | EP 1987-401018 | 19870504 |
| EP 245170 | B1 | 19891129 | | | |
| R: CH, DE, | GB, IT. | , Li, NL, | 5E | | |
| FR 2598421 | A1 | 19871113 | | FR 1986-6517 | 19860506 |
| FR 2598421 | B1 | 19880819 | | | |
| US 4912097 | Α | 19900327 | | US 1987-44958 | 19870430 |
| HU 44793 | A2 | 19880428 | | HU 1987-2007 | 19870505 |
| HU 196224 | В | 19881028 | | | |
| JP 62294694 | A2 | 19871222 | | JP 1987-109059 | 19870506 |
| DRITY APPLN. INFO. | : | | FR | 1986-6517 | 19860506 |
| ED COUDCE/C). | ~ > / | | . 1 2044 | ra | |

AZ 199/122 JF 199-19950

RITY APPLN. INFO.:

RR SOURCE(S): CASREACT 109:129463

Title steroids I [R1 = C2-8 alkynyl (un) substituted by OH, halo, trialkylsiyl, alkoxy, alkylthio, dialkylamino, or oxo: R2 = C1-3 alkyl; NB-rings = Q1-Q5: D-ring = Q6, Q7: R3, R4 = H, C1-4 alkyl; R5 = H, OH, acycloxy, (un) substituted C1-6 alkoxy; R6 = H, C1-8 alkyl, C7-15 aralkyl; R7, R8 = H, OH, etc., R7R8 = lactones and related groups; Y2 = CHZCHZ, CHICH, 1, 2-cyclopropanediyl, CHRSCH2, CHZCHR10: R9, R10 = C1-4 alkyl] are prepd. for use as progestogens, antiprogestogens, and/or antiglucocorticoids. 3,3-Ethylenedioxy-5,10-epoxy-estr-9[11]-en-17-one was treated with CH2:CHCHZMgBr and deprotected and dehydrated (NH4OH in aq. MeOH, then aq. HC1) to give (ethylnylphenyl) allylhydroxyestradienone II. At 10-OH in vitro, II gave 994 reversal of the dexamethasone-induced redn. of uridine uptake by rat thymocytes (S. times. 10-8M dexamethasone). Tablets were prepd. from 50 mg of the 17. alpha. (chlorothynyl) analog of III, and 120 mg of a mixt. of talc, starch, and Mg stearate.

116421-73-99 116421-74-OP

RL: SFN (Synthetic preparation); USES (Uses)

19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(1-propynyl)phenyl]-, (Ilbeta.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 28 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued)

96285-50-6 CAPLUS 18,19-Dinorpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-13-ethyl-, (11.bets.,13.alpha.)- (9CI) (CA INDEX NAME)

L5 ANSWER 29 OF 33 CAPLUS COPYRIGHT 2003 ACS

116421-74-0 CAPLUS 19-Norpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-(4-ethynylphenyl)-, (11.beta.)- (9CI) (CA INDEX NAME)

L5 ANSWER 30 0F 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 1987:5324 CAPLUS
106:5524 11.beta.-Phenylgonanes and pharmaceutical compositions containing them
Neef, Guenter: Weechert, Rudolf; Ottow, Eckard; Rohde, Ralph; Beier, Sybille: Elger, Walter; Henderson, David Schering A.-G., Fed. Rep. Ger.

DOCUMENT TYPE: EUR. PART ASPLOYED FROM A COUNT: 2

DOCUMENT TYPE: PART ASPLOYED FROM A COUNT: 2

CODEN: EXXDU

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|-------------------|------------|-----------|--------------------|----------|
| | | | | |
| EP 190759 | A2 | 19860813 | EP 1986-101548 | 19860206 |
| EP 190759 | A3 | 19861120 | | |
| EP 190759 | B1 | 19890830 | | |
| R: AT, | BE, CH, DE | , FR, GB, | IT, LI, LU, NL, SE | |
| DE 3504421 | A1 | 19860807 | DE 1985-3504421 | 19850207 |
| DE 3527517 | A1 | 19870129 | DE 1985-3527517 | 19850729 |
| AT 45956 | E | 19890915 | AT 1986-101548 | 19860206 |
| PRIORITY APPLN. I | NFO.: | | DE 1985-3504421 | 19850207 |
| | | | DE 1985-3527517 | 19850729 |
| | | | EP 1986-101548 | 19860206 |

DE 1985-3527517 19850729

DE 1985-3527517 19850729

EF 1986-101548 19860206

OTHER SOURCE(S): CASREACT 106:5327

AB 11.beta.-Phenylgonane derivs. I (2 = 0, CH2, bond; X = 0, NOH; R1 = 3- or 4-bydrocarbyl contg. C:X; R2 = .alpha.- or .beta.-Me or -Et; R3 and R4 = various group combinations (e.g. R3 or R4 = OH, acyloxy, other = (un) substituted C.tplbond.CH, R3R4 = CHICHIC202; R5-8 = H, OH, alkyl, alkoxy, acyloxy, halo] were prepd. as antigestagens and antiglucocorticoids, with a notable dissocn. of the two activities. Thus, 4-BrCGH4Ac was ketalized with Me2C(CH2OH)2, and the ketal was coupled with epoxyestrenol deriv. II by a CU-catalyzed Griganard reaction. The resulting arylgonane deriv. III (R3 = OH, R4 = H) was oxidized to give III (R3R4 = 0), which underwent alkynylation by Lic.tplbond.CMe or Lic.tplbond.CR9, R9 = Me or CH2OTHP). The former was hydrolyzed by aq. HOAc, and the latter was hydrogenated and then hydrolyzed by aq. HOAc, and the latter was hydrogenated and then hydrolyzed, to give IV (R4 = C.tplbond.CMe) (V) and (2)-IV (R4 = CR:CHCH2OH) (V1). V and VI showed, resp., 10- and 30-fold the abortifacient activity of the known compd. RU-38406 in gravid rats, while showing 301 and <11 of its antiglucocorticoid activity.

IT 10314-79-2 RL BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified), SFN (Synthetic preparation); BIOL (Biological study, PREP (Preparation) (prepn. of, as antigestagen and antiglucocorticoid)

RN 105114-79-2 CAFIUS

CN Benzaldehyde, 4-[(11.beta., 13.alpha.)-17-(acetyloxy)-3,20-dioxo-19-norpregna-4,9-dien-11-y1]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L5 ANSWER 31 OF 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 1987:5323 CAPLUS
DOCUMENT NUMBER: 1087:5323 CAPLUS
106:5323 CAPLUS
106:5323 CAPLUS
106:5323 CAPLUS
106:5322 CAPLUS
106:522 CAPLUS
106:522

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| PATENT NO. | KIND | DATE | | AP | PLICATION NO. | DATE |
|----------------------|------|----------|-------|----|---------------|----------|
| DE 3504421 | A1 | 19860807 | | DE | 1985-3504421 | 19850207 |
| AU 8652913 | A1 | 19860814 | | | 1986-52913 | 19860131 |
| AU 580843 | B2 | 19890202 | | | | |
| IL 77762 | A1 | 19920818 | | ΙL | 1986-77762 | 19860202 |
| CN 86100994 | A | 19861008 | | CN | 1986-100994 | 19860203 |
| CN 1033753 | В | 19970108 | | | | |
| ES 551625 | A1 | 19861216 | | ES | 1986-551625 | 19860204 |
| DK 8600560 | A | 19860808 | | DK | 1986-560 | 19860205 |
| DK 161709 | В | 19910805 | | | | |
| DK 161709 | С | 19920113 | | ٠, | | |
| NO 8600425 | A | 19860808 | | NO | 1986-425 | 19860206 |
| NO 171994 | В | 19930215 | | | | |
| NO 171994 | С | 19930526 | | | | |
| EP 190759 | A2 | 19860813 | | EP | 1986-101548 | 19860206 |
| EP 190759 | A3 | 19861120 | | | | |
| EP 190759 | B1 | 19890830 | | | | |
| | | | IT, L | | LU, NL, SE | |
| HU 40453 | A2 | 19861228 | | ΗU | 1986-499 | 19860206 |
| HU 194904 | В | 19880328 | | | | |
| DD 261166 | A5 | 19881019 | | | 1986-286860 | 19860206 |
| AT 45956 | E | 19890915 | | | 1986-101548 | 19860206 |
| CA 1310630 | A1 | 19921124 | | | 1986-501252 | 19860206 |
| FI 8600559 | A | 19860808 | | FΙ | 1986-559 | 19860207 |
| FI 85377 | В | 19911231 | | | | |
| FI 85377 | С | 19920410 | | | | |
| JP 61183296 | A2 | 19860815 | | J₽ | 1986-24260 | 19860207 |
| JP 04037080 | B4 | 19920618 | | | | |
| ZA 8600936 | A | 19860924 | | | 1986-936 | 19860207 |
| US 5089635 | A | 19920218 | | | 1986-827050 | 19860207 |
| NO 8604209 | A | 19860808 | | NO | 1986-4209 | 19861021 |
| NO 170285 | В | 19920622 | | | | |
| NO 170285 | С | 19920930 | | | | |
| RIORITY APPLN. INFO. | : | | | | 85-3504421 | 19850207 |
| | | | | | 85-3527517 | 19850729 |
| | | | | | 96-101549 | 19860206 |
| | | | | | | |

EP 1986-101548 19860206
NO 1986-425 19860206
Gonanes I [AB = 0, CH2, bond; X = 0, NOH: n = 0, 1; R1 = H, Cl-4 alkyl; R2 = Me, Et; R3, R4 = OR, acyloxy, alkynyl, acyl, Me, H, (substituted) alkyl, alkenyl, tetrahydrofuran-5-on-2-yl], useful as contraceptives, antiprogestins, and antiglucocorticoids (data given), were prepd. 17.alpha.-Ethynyl-11.beta.-(4-formylphenyl)-17.beta.-hydroxy-4,9-estradien-3-one was prepd. in 5 steps from 4-BrCGH4CNO, (HOCH2) 2CMe2, HC(OMe)3, and 4-MeCGH4SO3H.

RL: BAC (Biological activity or effector, except adverse); BSU (Biological

ANSWER 30 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued)

- ANSWER 31 OF 33 CAPLUS COPYRIGHT 2003 ACS (Continued) study, unclassified), SPN (Synthetic preparation), BIOL (Biological study); PREP (Preparation) (prepn. of, as antigestagen and antiglucocorticoid) 105114-79-2 CAPLUS Benzaldehyde, 4-[(11.beta.,13.alpha.)-17-(acetyloxy)-3,20-dioxo-19-norpregna-4,9-dien-11-y1]- (9CI) (CA INDEX NAME)

L5 ANSWER 32 OF 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 1986:34230 CAPLUS
DOCUMENT NUMBER: 104:34230 CAPLUS
TITLE: New steroids with antiprogestational and antiglucocorticoid activities
AUTHOR(S): Neef, Guenter Beier, Sybille: Elger, Walter, Henderson, David, Wiechert, Rudolf
CORPORATE SOURCE: Res. Lab., Schering A.-G./Bergkamen, Berlin, D-1000/65, Fed. Rep. Ger.
SOURCE: Steroids (1984), 44(4), 349-72
CODE: STEDAM; ISSN: 0039-128X
DOCUMENT TYPE: Journal
LANGUAGE: English
AB C-11 substituted 19-norsteroids I and II (R - HeO, F, MeZN; R1 = HO, AcO, HC. tplbond.C, HOCHZCHZCHZ; R2 = HO, Ac, HC. tplbond.C, HC. tplbond.C, HOCHZCHZCHZ; R2 = HO, Ac, HC. tplbond.C, HC. tplbond.C, HOCHZCHZCHZ; R2 = HO, Ac, HC. tplbond.C, HC. tplbond.C, HOCHZCHZCHZ; R2 = HO, Ac, HC. tplbond.C, HC

Absolute stereochemistry.

ANSWER 33 OF 33 CAPLUS COPYRIGHT 2003 ACS

96285-50-6 CAPLUS
18,19-Dinorpregna-4,9-diene-3,20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-13-ethyl-, (11.beta.,13.alpha.)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L5 ANSWER 33 OF 33 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 1985:406617 CAPLUS
DOCUMENT NUMBER: 103:6617
TITLE: 13.alpha.~Alkylgonanes and pharmaceutical compositions

INVENTOR(S):

13.alpha.-Alkylgonanes and pharmaceutical compose containing them
Neef, Guenter: Sauer, Gerhard; Wiechert, Rudolf; Beier, Sybille; Elger, Walter; Henderson, David; Rohde, Ralph
Schering A.-G., Fed. Rep. Ger.
Eur. Fat. Appl., 34 pp.
CODEN: EPXXDW
Patent
German
4

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE:

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|--------------------|-------------|----------|--------------------|----------|
| | | | | |
| EP 129499 | A2 | 19841227 | EP 1984-730062 | 19840613 |
| EP 129499 | A3 | 19851009 | | |
| EP 129499 | B1 | 19871209 | | |
| R: AT, I | BE, CH, DE, | FR, GB, | IT, LI, LU, NL, SE | |
| DE 3321826 | A1 | 19841220 | DE 1983-3321826 | 19830615 |
| DE 3413036 | A1 | 19851017 | DE 1984-3413036 | 19840404 |
| AT 31313 | E | 19871215 | AT 1984-730062 | 19840613 |
| PRIORITY APPLN. IN | NFO.: | | DE 1983-3321826 | 19830615 |
| | | | DE 1984-3413036 | 19840404 |
| | | | PR 1004 7300CD | 1001000 |

DE 1983-3321826 19830615

DE 1984-3430362 19840613

Phenylalkylgonenes I [R = H, alkyl; RI = amino, alkylamino, 5- or 6-membered heterocycle ring radical, alkoxy; R2 - H, Me, Et; R3 = alkyl, alkylatinylalkyl, alkoxyalkenyl, alkynyl, cyanoalkyl, Ac, HOHZCO; R4 = HO, alkoxy, acyloxy; R3R4 = 5-oxodihydrofuran-2(3H)-ylidene] were preped, via epimerization of estrene derives, and possessed antigestagenic and post-cottal contraceptive activities. Thus, the (aminophenyl)estrenone ketal II was photolyzed in THF using a Hg high-pressure lamp to give the C-13 epimer of II, which underwent successive addn. ceaction with LiC.tplbond.CCH20-THP (THP = tetrahydropyranyl), hydrogenation, and hydrolysis to give the (hydroxypropyl)gonadiene III. At 10 mg/animal/day III had a 1004 abortion rate in rats.

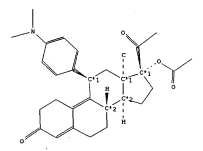
96285-40-4P 96285-50-6P
RL: SPN (Synthetic preparation); PREP (Preparation)
(prepn. of)
96285-40-4 CAPLUS
19-Norpregna-4, 9-diene-3, 20-dione, 17-(acetyloxy)-11-[4-(dimethylamino)phenyl]-, (11.beta.,13.alpha.)- (9CI) (CA INDEX NAME)

09/526,855 Page 31

=> d all

ANSWER 1 OF 1 BEILSTEIN COPYRIGHT 2003 BEILSTEIN CDS MDL

5673666
96285-40-4, 126784-99-4
17. alpha.-acetoxy-11.beta.-(4dimethylaminophenyl)-13.alpha.-methyl18,19-dinor-pregna-4,9-dinen-3,20-dione
acetic acid 17-acetyl-11-(4-dimethylaminophenyl)-13-methyl-3-oxo2,3,6,7,8,11,12,13,14,15,16,17-dodecahydro1H-cyclopenta<a>phenanthren-17-yl ester
C30 H37 N O4
475.63
15934, 2817, 1155
Stereo compound
isocyclic
5000625
5427628
6-14
1933/02/12 Beilstein Records (BRN): CAS Reg. No. (RN): Chemical Name (CN): 5673666 Autonom Name (AUN): Molec. Formula (MF):
Molecular Weight (MW):
Lawson Number (LM):
File Segment (FS):
Compound Type (CTYPE):
Constitution ID (CONSID):
Tautomer ID (TAUTIO):
Beilstein Citation (BSO):
Entry Date (DED):
Update Date (DUPD):



Atom/Bond Notes: CIP Descriptor: R
 CIP Descriptor: S

Field Availability:

| Code | Name | Occurrence |
|------|---------------------|------------|
| | | |
| BRN | Beilstein Records | 1 |
| RN | CAS Registry Number | 2 |

ANSWER 1 OF 1 BEILSTEIN COPYRIGHT 2003 BEILSTEIN CDS MDL (Continue Nucleus (.NUC): 1H
Solvents (.SOL): CDC13
Reference(s): 1. Neef, Guenter; Beier, Sybille; Elger, Walter; Henderson, David; Wiechert, Rudolf, Steroids, CODEN: STEDAM, 44(4), <1984>, 349-372; BABS-5685203

Bands | KBr 1 1 1 1

Reference(s):
1. Neef, Guenter: Beier, Sybille: Elger, Walter: Henderson, David: Wiechert, Rudolf, Steroids, CODEN: STEDAM, 44(4), <1984>, 349-372: BABS-5685283

Notes(s): 1. 1736 - 1612 cm**(-1)

Pharmacological Data: PHARM Note(s) (.COM):

reversal of dexamethasone induced tyrosine aminotransferase activity in rat hepatoma cells (antiglucocorticoid activity)

Reference(s):

1. Neef, Guenter; Beier, Sybille; Elger, Walter; Henderson, David; Wiechert, Rudolf, Steroids, CODEN: STEDAM, 44(4), <1984>, 349-372; BABS-5685283

Reaction ID (.ID): Reactant BRN (.RBRN): Reactant (.RCT):

2373868
5657948, 385737
11.beta.-(4-dimethylaminophenyl)-17.alpha.-hydroxy-13.alpha.-methyl-18,19-dinor-pregna-4,9-diene-3,20-dione, acetic acid anhydride
5673666
17.alpha.-acetoxy-11.beta.-(4-dimethylaminophenyl)-13.alpha.-methyl-18,19-dinor-pregna-4,9-diene-3,20-dione

Product BRN (.PBRN): Product (.PRO):

No. of React. Details (.NVAR):

Reaction Details:

Reaction RID (.RID): 2373868.1

Reaction Classification (.CL): Preparation
Yield (.YDT): 93 percent (BRN=5673666)
Reagent (.RGT): 4-dimethylaminopyridine
Solvent (.SOL): toluene
Time (.TIM): 14 hour(s)
Other Conditions (.COND): Ambient temperature
Reference(s):

1. Neef, Guenter; Beier, Sybille; Elger, Walter; Henderson, David;
Wiechert, Rudolf, Steroids, CODEN: STEDAM, 44(4), <1984>, 349-372;

ANSWER 1 OF 1 BEILSTEIN COPYRIGHT 2003 BEILSTEIN CDS MDL
CN Chemical Name 1
AUN Autonomname 1
NF Molecular Formula 1
FW Formular Weight 1
LN Lawson Number 3
FS File Segment 1
CTYPE Compound Type 1
CONSID Constitution ID 1
TAUTID 1
BSO Beilstein Citation 1
BD Entry Date 1
LD Letter Compound Type 1
LTAUTID 1
LTAUTID

This substance also occurs in Reaction Documents:

| Code | Name | Occurrenc |
|-------|-------------------------------|-----------|
| RX | Reaction Documents | |
| RXPRO | Substance is Reaction Product | |

Melting Point: Value (MP) (Cel) |Solvent |(.SOL) |Ref. 194 - 195 jethyl acetate, hexanei 1

Reference(s):

Neef, Guenter; Beier, Sybille; Elger, Walter; Henderson, David; Wiechert, Rudolf, Steroids, CODEN: STEDAM, 44(4), <1984>, 349-372; BABS-5685283

Optical Rotatory Power:
Value |Type |Concentr.
(ORP) |(.TYP) |(.C)
(deg) | |Solvent | Wavelen. |(.SOL) | (.W) | | (nm) | Temp. | | (.T) | | (Cel) | 372.3 1 589 25 [[alpha] | 10.39 g/100ml|CHC13

Reference (s):

Rudolf, Steroids, CODEN: STEDAM, 44(4), <1984>, 349-372; BABS-5685283

Nuclear Magnetic Resonance:

Description (.KW):

Chemical shifts

ANSWER 1 OF 1 BEILSTEIN COPYRIGHT 2003 BEILSTEIN CDS MDL (Continued) BABS-5685283

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L5

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L1 STRUCTURE UPLOADED

L2 6 S L1

L3 62 S L1 FULL

FILE 'USPATFULL' ENTERED AT 15:51:54 ON 28 APR 2003

L4 11 S L3

FILE 'CAPLUS' ENTERED AT 15:53:08 ON 28 APR 2003 33 S L3

FILE 'BEILSTEIN' ENTERED AT 15:55:48 ON 28 APR 2003 L6 1 S L3

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